

STIC Search Report Biotech-Chem Library

STIC Database Tracking Number: 97806

TO: Jennifer Kim

Location: 2d17 / 2b19 Tuesday, July 01, 2003

Art Unit: 1617 Phone: 308-2232

Serial Number: 09 / 719770

From: Jan Delaval

Location: Biotech-Chem Library

CM1-1E07

Phone: 308-4498

jan.delaval@uspto.gov

Search Notes

Jan Delaval Reference Librarian Biotechnology & Chemical Library CM1 1E07 – 703-308-4498 jan.delaval@uspto.gov



Fan Deleval

Access DB# 97806

SEARCH REQUEST FORM

Scientific and Technical Information Center

Requester's Full Name: Jenniter Kin Examiner #: 19469 Date: 1/1/03
Art Unit: /6/1 Phone Number 30 & -2232 Serial Number: 109 /119110
Mail Box and Bldg/Room Location: 20/1 Results Format Preferred (circle): RAPER DISK E-MAIL
If more than one search is submitted, please prioritize searches in order of need.
Please provide a detailed statement of the search topic, and describe as specifically as possible the subject matter to be searched.
Include the elected species or structures, keywords, synonyms, acronyms, and registry numbers, and combine with the concept or utility of the invention. Define any terms that may have a special meaning. Give examples or relevant citations, authors, etc, if
known. Please attach a copy of the cover sheet, pertinent claims, and abstract.
William for treating diseases medicales sof
Known. Please attach a copy of the cover sheet, pertinent claims, and abstract. Title of Invention: Mothods + Compositions for freating descrees modicated by Inventors (please provide full names): Laurence (form ma)
Inventors (please provide full names): Lawrence (tom man)
Earliest Priority Filing Date: 6/19/98
For Sequence Searches Only Please include all pertinent information (parent, child, divisional, or issued patent numbers) along with the appropriate serial number.
) Please search claims 1-3, 15+16 based on the
disease tisted in claim 3 with active agent
disease tisted in claim 3 with
of claim 8 of Monodansyl cadaverine,
e) Please provide registry# &
) Frust 1.
Jan Delaval THX Reference Librarian
Biotechnology & Chemical Library
CM1 1E07 – 703-308-4498 [77] jan.delaval@uspto.gov

COTA TOTA TICITA AND TO THE COLUMN TO THE CO



STIC SEARCH RESULTS

Biotech-Chem Library

Questions about the scope or the results of the search? Contact the searcher or contact:

Mary Hale, Information Branch Supervisor 308-4258, CM1-1E01

Voluntary Results Feedback Form							
> I am an examiner in Workgroup: Example: 1610							
Relevant prior art found, search results used as follows:							
☐ 102 rejection							
☐ 103 rejection							
Cited as being of interest.							
Helped examiner better understand the invention.							
Helped examiner better understand the state of the art in their technology.							
Types of relevant prior art found:							
☐ Foreign Patent(s)							
 Non-Patent Literature (journal articles, conference proceedings, new product announcements etc.) 							
> Relevant prior art not found:							
Results verified the lack of relevant prior art (helped determine patentability).							
Results were not useful in determining patentability or understanding the invention.							
Comments:							

Drop off or send completed forms to STIC/Biotech-Chem Library CM1 = Circ Desk



=> fil hcaplus FILE 'HCAPLUS' ENTERED AT 16:07:58 ON 01 JUL 2003 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2003 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 1 Jul 2003 VOL 139 ISS 1 FILE LAST UPDATED: 30 Jun 2003 (20030630/ED)

This file contains CAS Registry Numbers for easy and accurate | Jan Delaval substance identification. | Reference Librarian

Biotechnology & Chemical Library CMI 1E07 – 703-308-4498 jan.delaval@uspta.gov

=> d all hitstr tot

L71 ANSWER 1 OF 11 HCAPLUS COPYRIGHT 2003 ACS

AN 2002:224182 HCAPLUS

- TI Prolonged survival and decreased abnormal movements in transgenic model of Huntington disease, with administration of the transglutaminase Inhibitor cystamine. [Erratum to document cited in CA137:592]
- AU Karpuj, Marcela V.; Becher, Mark W.; Springer, Joe E.; Chabas, Dorothee; Youssef, Sawsan; Pedotti, Rosetta; Mithcell, Dennis; Steinman, Lawrence
- CS Department of Neurological Sciences, Stanford University, Stanford, CA, USA
- SO Nature Medicine (New York, NY, United States) (2002), 8(3), 303 CODEN: NAMEFI; ISSN: 1078-8956
- PB Nature America Inc.
- DT Journal; Errata
- LA English
- CC 1-11 (Pharmacology)
- AB An erratum.
- ST erratum transglutaminase inhibitor cystamine Huntington disease huntingtin polyglutamine; transglutaminase inhibitor cystamine Huntington disease huntingtin polyglutamine erratum
- IT INDEXING IN PROGRESS
- IT Nervous system

(Huntington's chorea; transglutaminase

inhibitor cystamine effect on survival and abnormal movements in a transgenic model of **Huntington** disease (Erratum))

IT Gene, animal

RL: BSU (Biological study, unclassified); BIOL (Biological study) (dnaj; transglutaminase inhibitor cystamine effect on survival and abnormal movements in a transgenic model of Huntington disease in relation to transcription of (Erratum))

IT Proteins

17°

RL: BSU (Biological study, unclassified); BIOL (Biological study) (huntingtin; transglutaminase inhibitor cystamine effect on survival and abnormal movements in a transgenic model of Huntington disease involving exon 1 of huntingtin contg. an expanded polyglutamine repeat (Erratum))

```
51-85-4, Cystamine
     RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (transglutaminase inhibitor cystamine effect on survival and
        abnormal movements in a transgenic model of Huntington
        disease (Erratum))
IT
     INDEXING IN PROGRESS
     51-85-4, Cystamine
ΙT
     RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (transglutaminase inhibitor cystamine effect on survival and
        abnormal movements in a transgenic model of Huntington
        disease (Erratum))
     51-85-4 HCAPLUS
RN
     Ethanamine, 2,2'-dithiobis- (9CI)
                                        (CA INDEX NAME)
CN
H2N-CH2-CH2-S-S-CH2-CH2-NH2
     ANSWER 2 OF 11 HCAPLUS COPYRIGHT 2003 ACS
L71
     2002:123990 HCAPLUS
AN
DN
     137:592
     Prolonged survival and decreased abnormal movements in transgenic model of
ΤI
     Huntington disease, with administration of the
     transglutaminase inhibitor cystamine
     Karpuj, Marcela V.; Becher, Mark W.; Springer, Joe E.; Chabas, Dorothee;
ΑU
     Youssef, Sawsan; Pedotti, Rosetta; Mitchell, Dennis; Steinman,
     Lawrence
     Department of Neurological Sciences, Stanford University, Stanford, CA,
CS
     USA
     Nature Medicine (New York, NY, United States) (2)(2), 8(2), 143-149
SO
     CODEN: NAMEFI; ISSN: 1078-8956
PB
     Nature America Inc.
     Journal
DΤ
LA
     English
CC
     1-11 (Pharmacology)
     An expanded polyglutamine domain in huntingtin underlies the pathogenic
AB
     events in Huntington disease (HD), characterized by
     chorea, dementia and severe wt. loss, culminating in death.
     Transglutaminase (TGase) may be crit. in the pathogenesis; via
     crosslinking huntingtin. Administration of the TGase competitive
     inhibitor cystamine to transgenic mice expressing exon 1 of huntingtin
     contq. an expanded polyglutamine repeat, altered the course of their
     HD-like disease. Cystamine given i.p. entered the brain, where it
     inhibited TGase activity. When treatment was begun after the appearance
     of abnormal movements, cystamine extended survival, reduced the assocd.
     tremor and abnormal movements and ameliorated wt. loss. Treatment did not
     influence the appearance or frequency of neuronal nuclear inclusions.
     Unexpectedly, cystamine increased transcription of one of the two genes
     shown to be neuroprotective for polyglutamine toxicity in Drosophila, dnaj
     (also known as HD/1 and Hsp40 in humans and mice, resp.). Inhibition of
     TGase provides a new treatment strategy for HD and other polyglutamine
     diseases.
     transglutaminase inhibitor cystamine Huntington
ST
     disease huntingtin polyglutamine
TΤ
     Nervous system, disease
        (Huntington's chorea; transglutaminase
        inhibitor cystamine effect on survival and abnormal movements in a
        transgenic model of Huntington disease)
```

RL: BSU (Biological study, unclassified); BIOL (Biological study)

IT

Gene, animal

(dnaj; transglutaminase inhibitor cystamine effect on survival and abnormal movements in a transgenic model of Huntington disease in relation to transcription of)

IT Proteins

RL: BSU (Biological study, unclassified); BIOL (Biological study) (huntingtin; transglutaminase inhibitor cystamine effect on survival and abnormal movements in a transgenic model of Huntington disease involving exon 1 of huntingtin contg. an expanded polyglutamine repeat)

IT 80146-85-6, Transglutaminase

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(inhibitors; transglutaminase inhibitor cystamine effect on

(inhibitors; transglutaminase inhibitor cystamine effect on survival and abnormal movements in a transgenic model of Huntington disease)

IT **51-85-4**, Cystamine

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(transglutaminase inhibitor cystamine effect on survival and abnormal movements in a transgenic model of **Huntington** disease)

IT 26700-71-0, Polyglutamine 69864-43-3, Polyglutamine

RL: BSU (Biological study, unclassified); BIOL (Biological study)
(transglutaminase inhibitor cystamine effect on survival and
abnormal movements in a transgenic model of Huntington
disease involving exon 1 of huntingtin contg. an expanded polyglutamine repeat)

RE.CNT 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD RE

- (1) Bates, G; Brain Pathol 1998, V8, P699 HCAPLUS
- (2) Becher, M; Neurobiol Disease 1998, V4, P387 HCAPLUS
- (3) Chabas, D; Science 2001, V294, P1731 HCAPLUS
- (4) Chen, M; Nature Med 2000, V6, P797 HCAPLUS
- (5) Clarke, G; Nature 2000, V406, P195 HCAPLUS
- (6) Cooper, A; J Neurochem 1997, V69, P431 HCAPLUS
- (7) Cummings, C; Nature Genet 1998, V19, P148 HCAPLUS
- (8) Curtis, C; Methods Enzymol 1976, V45, P177 HCAPLUS
- (9) Davies, S; Cell 1997, V90, P537 HCAPLUS
- (10) DiFiglia, M; Science 1997, V277, P1990 HCAPLUS
- (11) Ferrante, R; J Neurosci 2000, V20, P4389 HCAPLUS
- (12) Folk, J; Ann Rev Biochem 1980, V49, P517 HCAPLUS
- (13) Folk, J; Biochim Biophys Acta 1966, V122, P244 HCAPLUS
- (14) Green, H; Cell 1993, V74, P955 HCAPLUS
- (15) Huang, C; Somatic Cell Mol Genet 1998, V24, P217 HCAPLUS
- (16) Huntington's Disease Collaborative Research Group; Cell 1993, V72, P971 HCAPLUS
- (17) Igarashi, S; Nature Genet 1998, V18, P111 HCAPLUS
- (18) Isupov, M; Structure 1996, V4, P801 HCAPLUS
- (19) Jeitner, T; J Neurochem 2001, V79, P1109 HCAPLUS
- (20) Kahlem, P; Mol Cell 1998, V1, P595 HCAPLUS
- (21) Karpuj, M; Proc Natl Acad Sci 1999, V96, P7388 HCAPLUS
- (22) Kazemi Esfarjani, P; Science 2000, V287, P1837 HCAPLUS
- (23) Kobayashi, Y; J Biol Chem 2000, V275, P8772 HCAPLUS
- (24) Lesort, M; J Neurochem 1999, V73, P2018 HCAPLUS
- (25) Lorand, L; Biochemistry 1979, V18, P1756 HCAPLUS
- (26) Lorand, L; Mol Cell Biochem 1984, V58, P25
- (27) Lorand, L; Nature Genet 1998, V20, P231 HCAPLUS
- (28) Lorand, L; Proc Natl Acad Sci 1996, V93, P14310 HCAPLUS
- (29) Lorand, L; Proc Natl Acad Sci 1997, V93, P14310
- (30) Luthi-Carter, R; Hum Mol Genet 2000, V9, P1259 HCAPLUS
- (31) Mangiarini, L; Cell 1996, V87, P493 HCAPLUS
- (32) Molberg, O; Nature Med 1998, V4, P713 HCAPLUS
- (33) Newcomb, R; J Biol Chem 1997, V272, P11276 HCAPLUS

```
(34) Ona, V; Nature 1999, V399, P263 HCAPLUS
(35) Ordway, J; Cell 1997, V91, P753 HCAPLUS
(36) Orr, H; Cell 2000, V101, P1 HCAPLUS
(37) Perutz, M; Trends Biochem Sci 1999, V24, P58 HCAPLUS
(38) Rittling, S; Biochem Biophys Res Commun 1998, V250, P287 HCAPLUS
(39) Sathasivam, K; Hum Mol Genet 1999, V8, P813 HCAPLUS
(40) Saudou, P; Cell 1998, V95, P55
(41) Scheufler, C; Cell 2000, V101, P199 HCAPLUS
(42) Sisodia, S; Cell 1998, V95, P1 HCAPLUS
(43) Springer, J; Nature Med 1999, V5, P943 HCAPLUS
(44) Voehringer, D; Proc Natl Acad Sci 2000, V97, P2680 HCAPLUS
(45) Yamamoto, A; Cell 2000, V101, P57 HCAPLUS
(46) Yu, S; Science 1999, V284, P336 HCAPLUS
(47) Zainelli, G; Soc Neurosci Absracts 2000, V26, P1297
TΤ
     80146-85-6, Transglutaminase
     RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (inhibitors; transglutaminase inhibitor cystamine effect on
        survival and abnormal movements in a transgenic model of
        Huntington disease)
RN
     80146-85-6 HCAPLUS
CN
     Glutamyltransferase, glutaminylpeptide .gamma.- (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     51-85-4, Cystamine
     RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (transglutaminase inhibitor cystamine effect on survival and
        abnormal movements in a transgenic model of Huntington
        disease)
RN
     51-85-4 HCAPLUS
CN
     Ethanamine, 2,2'-dithiobis- (9CI)
                                        (CA INDEX NAME)
H2N-CH2-CH2-S-S-CH2-CH2-NH2
IT
     26700-71-0, Polyglutamine 69864-43-3, Polyglutamine
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (transglutaminase inhibitor cystamine effect on survival and
        abnormal movements in a transgenic model of Huntington
        disease involving exon 1 of huntingtin contg. an expanded polyglutamine
        repeat)
RN
     26700-71-0
                HCAPLUS
                                      (CA INDEX NAME)
CN
     L-Glutamine, homopolymer (9CI)
     CM
          1
     CRN
          56-85-9
     CMF
         C5 H10 N2 O3
Absolute stereochemistry.
              NH<sub>2</sub>
```

Poly[imino[(1S)-1-(3-amino-3-oxopropyl)-2-oxo-1,2-ethanediyl]] (9CI)

69864-43-3 HCAPLUS

RN

CN

INDEX NAME)

```
CH<sub>2</sub>-CH<sub>2</sub>-C-NH<sub>2</sub>

CH<sub>2</sub>-CH<sub>2</sub>-C-NH<sub>2</sub>

NH-CH-C----

O
II
O
In
```

```
ANSWER 3 OF 11 HCAPLUS COPYRIGHT 2003 ACS
L71
ΑN
     1999:811104
                 HCAPLUS
DN
     132:45002
     Methods and compositions for treating diseases mediated by
ΤI
     transglutaminase activity
     Steinman, Lawrence; Karpuj, Marcella V.
IN
     Yeda Research and Development Co. Ltd., Israel
PA
     PCT Int. Appl., 61 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
Τ<sub>τ</sub>Α
IC
     ICM A61K038-48
     ICS A61K031-13
CC
     1-12 (Pharmacology)
     Section cross-reference(s): 6/3
FAN.CNT 1
     PATENT NO.
                             DATE
                                             APPLICATION NO.
                       KIND
     WO 9965516
                             19991223
                                             WO 1999-US13615
                                                               19990617 <--
PΙ
                        Α1
            AE, AL, AM, AT, AÚ, XZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             DE, DK, EE, ES, FI, GR, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK,
             MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ,
             TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ,
             MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
                      GA, GN, GW, ML, MR, NE, SN, TD, TG
             CI, CM,
                        A1 : 20000105
                                             AU 1999-48239
                                                               19990617 <--
     AU 9948239
PRAI US 1998-89603P
                             19980617
                        Р
     WO 1999-US13615
                        Ŵ
                             19990617
                                        <--
AB
     Diseases mediated by transglutaminase, e.g. Huntington
     's Disease, spinobulbar atrophy,
     spinocerebellar ataxia, and
     dentatorubralpallidoluysian atrophy, as well as
     inflammatory diseases of the central nervous system, including multiple
     sclerosis, rheumatoid arthritis, and insulin-dependent diabetes mellitus,
     can be treated by administering a transglutaminase inhibitor,
     e.g. monodansyl cadaverine, monoamines and diamines
     such as cystamine or putrescine, etc.
ST
     transglutaminase inhibitor therapeutic; nervous system disease
     transglutaminase inhibitor; antiinflammatory antidiabetic
     transglutaminase inhibitor
IT
     Nervous system
         (Huntington's chorea; transglutaminase
        inhibitor-based methods and compns. for treating diseases mediated by
        transglutaminase activity)
IT
     Virus vectors
```

(and transkaryotic implantation; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) IT Disease, animal (atrophy, spinobulbar; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) ΙT Encephalomyelitis (autoimmune; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) ΙT Erythrocyte Erythrocyte (cell membrane, liposome hybrid; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transqlutaminase activity) Autoimmune disease TΤ (cell-mediated; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) IT Nervous system (central, inflammation; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) ΙT Brain (cerebellum; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) IT Brain (cortex; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) IT. Brain (corticular nuclear ext.; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) IT Nervous system (degeneration; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) ТТ Brain, disease (dentatorubral-pallidoluysian atrophy; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) Cell membrane IT Cell membrane (erythrocyte, liposome hybrid; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) IT Proteins, specific or class RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (huntingtin; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) Drug delivery systems IT (immunoliposomes; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) Diabetes mellitus IT (insulin-dependent; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity)

IT

Drug delivery systems

252868-74-9, 3: PN:

(liposomes; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) ΙT Aggregation (of polyQ-contg. proteins; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) ΙT Receptors RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (receptor-mediated gene delivery; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) ΙT Nervous system (spinocerebellar ataxia; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) IT Multiple sclerosis (therapeutic agents; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) Anti-inflammatory agents ΙT Antidiabetic agents Antirheumatic agents Drug delivery systems Gene therapy Lymphoblast Nervous system agents Retroviral vectors (transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) ΙT Antisense DNA RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (transqlutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) ITGene, animal RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (transglutaminase; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) 24991-23-9 25513-46-6, Polyglutamic acid IT RL: BSU (Biological study, unclassified); BIOL (Biological study) (polyQ-contg. protein aggregation; transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) 51-85-4, Cystamine 64-77-7, Tolbutamide 110-60-1 Putrescine 150-13-0 616-34-2, Glycine methyl ester 7758-98-7, Cupric sulfate, biological studies 10121-91-2 , Monodansyl cadaverine 74389-76-7 RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity) 80146-85-6, Transglutaminase ΙT RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity)

252868-73-8, 2: PN: WO9965516 SEQID: 3 unclaimed DNA

IT

WO9965516 SEQID: 4 unclaimed DNA

RL: PRP (Properties)

(unclaimed nucleotide sequence; methods and compns. for treating diseases mediated by transqlutaminase activity)

IT 252874-67-2

RL: PRP (Properties)

(unclaimed protein sequence; methods and compns. for treating diseases mediated by transglutaminase activity)

IT 252769-79-2

RL: PRP (Properties)

(unclaimed sequence; methods and compns. for treating diseases mediated by transglutaminase activity)

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD RE

(1) Heska Corporation USA; WO 9824887 A2 1998 HCAPLUS

(2) O'Hara; US 5514579 A 1996 HCAPLUS

(3) Victoria University of Manchester; WO 9804245 A1 1998 HCAPLUS

IT 24991-23-9 25513-46-6, Polyglutamic acid

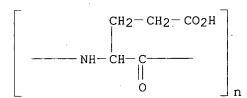
RL: BSU (Biological study, unclassified); BIOL (Biological study)

(polyQ-contg. protein aggregation; transglutaminase

inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity)

RN 24991-23-9 HCAPLUS

CN Poly[imino[(1S)-1-(2-carboxyethyl)-2-oxo-1,2-ethanediyl]] (9CI) (CA INDEX NAME)



RN 25513-46-6 HCAPLUS

CN L-Glutamic acid, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 56-86-0

CMF C5 H9 N O4

Absolute stereochemistry.

IT 51-85-4, Cystamine 64-77-7, Tolbutamide 110-60-1

, Putrescine 150-13-0 616-34-2, Glycine methyl ester

7758-98-7, Cupric sulfate, biological studies 10121-91-2

, Monodansyl cadaverine 74389-76-7

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity)

RN 51-85-4 HCAPLUS

CN Ethanamine, 2,2'-dithiobis- (9CI) (CA INDEX NAME)

 $H_2N-CH_2-CH_2-S-S-CH_2-CH_2-NH_2$

RN 64-77-7 HCAPLUS

CN Benzenesulfonamide, N-[(butylamino)carbonyl]-4-methyl- (9CI) (CA INDEX NAME)

RN 110-60-1 HCAPLUS

CN 1,4-Butanediamine (8CI, 9CI) (CA INDEX NAME)

 $H_2N-(CH_2)_4-NH_2$

RN 150-13-0 HCAPLUS

CN Benzoic acid, 4-amino- (9CI) (CA INDEX NAME)

RN 616-34-2 HCAPLUS

CN Glycine, methyl ester (6CI, 8CI, 9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{O} \\ || \\ \text{MeO-C-CH}_2\text{--NH}_2 \end{array}$$

RN 7758-98-7 HCAPLUS

CN Sulfuric acid copper(2+) salt (1:1) (8CI, 9CI) (CA INDEX NAME)

Cu(II)

RN 10121-91-2 HCAPLUS

CN 1-Naphthalenesulfonamide, N-(5-aminopentyl)-5-(dimethylamino)- (8CI, 9CI)

(CA INDEX NAME)

$$O = S - NH - (CH2)5 - NH2$$

$$NMe2$$

RN 74389-76-7 HCAPLUS

CN Norvaline, 5-diazo-4-oxo-N-[(phenylmethoxy)carbonyl]-, 4-nitrophenyl ester (9CI) (CA INDEX NAME)

IT 80146-85-6, Transglutaminase

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(transglutaminase inhibitor-based methods and compns. for treating diseases mediated by transglutaminase activity)

RN 80146-85-6 HCAPLUS

CN Glutamyltransferase, glutaminylpeptide .gamma.- (9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

L71 ANSWER 4 OF 11 HCAPLUS COPYRIGHT 2003 ACS

AN 1999:481278 HCAPLUS

DN 131:125479

TI Therapeutic agents for CAG repeat expansion disease

IN Tsuji, Shoji

PA Niigata University, Japan

SO Jpn. Kokai Tokkyo Koho, 19 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

IC ICM A61K045-00

ICS A61K045-00; A61K031-195; A61K038-00

CC · 1-12 (Pharmacology)

FAN CNT 1

FAN.CNT 1							
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
ΡI	JP 11209304	A2	19990803	JP 1998-27739	199/80126 <		
	JP 3012923	B2	20000228	•	\ /		
	AU 9913191	A1	19990812	AU 1999-13191	19 990122 <		
	US < 6355690	В1	20020312	US 1999-236002	1/9990122 <		
	EP 950406	A2	19991020	EP 1999-101063	19990125 <		
	EP 950406	A3	20001129				

ď.

```
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
                                           CA 1999-2260311 19990125 <--
    CA 2260311
                       С
                            20021217
PRAI JP 1998-27739
                            19980126
                                     <--
    Therapeutic agents for CAG repeat expansion disease comprise
    transglutaminase inhibitors i.e. cysteamine and monodansyl
    cadaverine. CAG repeat expansion disease is spinal and bulbar
    muscular atrophy, Huntington's disease,
    spinocerebeller ataxia type 2, hereditary dentatorubral
    pallidoluysian atrophy, Machado-Joseph disease or
    autosomal dominal cerebellar ataxia.
    CAG repeat expansion disease transglutaminase inhibitor;
ST
    cysteamine CAG repeat expansion disease; monodansyl
    cadaverine CAG repeat expansion disease
    Disease, animal
ΙT
        (CAG repeat expansion; therapeutic agents for CAG repeat expansion
       disease)
ΙT
    Nervous system
        (Huntington's chorea; therapeutic agents for CAG
       repeat expansion disease)
ΙT
    Nervous system
        (Machado-Joseph disease; therapeutic agents for CAG repeat expansion
       disease)
ΙT
    Nervous system
        (ataxia, spinocerebeller or autosomal dominal cerebellar;
        therapeutic agents for CAG repeat expansion disease)
ΙT
    Disease, animal
        (atrophy, hereditary dentatorubral
       pallidoluysian; therapeutic agents for CAG repeat expansion
       disease)
IT
    Spinal muscular atrophy
        (spinal and bulbar muscular atrophy; therapeutic agents for
       CAG repeat expansion disease)
IT
    80146-85-6, Transglutaminase
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
    study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES
     (Uses)
        (inhibitors; therapeutic agents for CAG repeat expansion disease)
    60-23-1, Cysteamine 10121-91-2, Monodansyl
ΤТ
    cadaverine
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
    study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES
     (Uses)
        (therapeutic agents for CAG repeat expansion disease)
    80146-85-6, Transglutaminase
TT
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
    study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES
     (Uses)
        (inhibitors; therapeutic agents for CAG repeat expansion disease)
RN
    80146-85-6 HCAPLUS
    Glutamyltransferase, glutaminylpeptide .gamma.- (9CI) (CA INDEX NAME)
CN
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
    10121-91-2, Monodansyl cadaverine
IT
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
    study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES
     (Uses)
        (therapeutic agents for CAG repeat expansion disease)
    10121-91-2 HCAPLUS
RN
    1-Naphthalenesulfonamide, N-(5-aminopentyl)-5-(dimethylamino)- (8CI, 9CI)
CN
     (CA INDEX NAME)
```

```
ANSWER 5 OF 11 HCAPLUS COPYRIGHT 2003 ACS
L71
     1998:406389 HCAPLUS
ΑN
     129:79862
DN
ΤI
     CAG repeat diseases and neuronal cell death
     Igarashi, Shuichi; Koide, Reiji; Shimohata, Takayoshi; Tsuji, Shoji
ΑU
CS
     Brain Res. Inst., Niigata Univ., Niigata, 951, Japan
     Jikken Igaku (1998), 16(10), 1277-1280
SO
     CODEN: JIIGEF; ISSN: 0288-5514
PB
     Yodosha
DT
     Journal; General Review
LA
     Japanese
     14-0 (Mammalian Pathological Biochemistry)
CC
     Section cross-reference(s): 3
     A review with 10 refs., on involvement of aggregates of proteins contg.
AΒ
     polyglutamine in mechanisms of CAG repeat diseases (Huntington's disease,
     Machado-Joseph disease, dentatorubral-pallidoluysian atrophy,
     etc.) and neuronal cell death. Involvement of transglutaminase
     in aggregate formation is also discussed.
     review CAG repeat disease polyglutamine aggregation; neuronal cell death
ST
     polyglutamine aggregation review; transglutaminase polyglutamine
     aggregation neuronal cell review
ΙT
     Nerve, disease
        (death; involvement of polyglutamine aggregation in CAG repeat diseases
        and neuronal cell death)
     Nervous system
TT
        (degeneration; involvement of polyglutamine aggregation in CAG repeat
        diseases and neuronal cell death)
ΙT
     Mutation
        (expansion; involvement of polyglutamine aggregation in CAG repeat
        diseases and neuronal cell death)
TΤ
     Repetitive DNA
     RL: ADV (Adverse effect, including toxicity); BIOL (Biological study)
        (involvement of polyglutamine aggregation in CAG repeat diseases and
        neuronal cell death)
     Gene, animal
IT
     RL: ADV (Adverse effect, including toxicity); PRP (Properties); BIOL
     (Biological study)
        (involvement of polyglutamine aggregation in CAG repeat diseases and
        neuronal cell death)
ΤT
     Cell death
        (neuron; involvement of polyglutamine aggregation in CAG repeat
```

RL: ADV (Adverse effect, including toxicity); BIOL (Biological study)

BPR (Biological process); BSU (Biological study, unclassified); BIOL

(involvement of polyglutamine aggregation in CAG repeat diseases and

RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence);

diseases and neuronal cell death)

26700-71-0, Polyglutamine **69864-43-3**, Polyglutamine

(Biological study); OCCU (Occurrence); PROC (Process)

80146-85-6, Transglutaminase

neuronal cell death)

IT

IT

(involvement of polyglutamine aggregation in CAG repeat diseases and neuronal cell death)

IT 101985-79-9, DCAG

RL: ADV (Adverse effect, including toxicity); BIOL (Biological study) (repeat; involvement of polyglutamine aggregation in CAG repeat diseases and neuronal cell death)

IT 80146-85-6, Transglutaminase

RL: ADV (Adverse effect, including toxicity); BIOL (Biological study) (involvement of polyglutamine aggregation in CAG repeat diseases and neuronal cell death)

RN 80146-85-6 HCAPLUS

CN Glutamyltransferase, glutaminylpeptide .gamma.- (9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

26700-71-0, Polyglutamine 69864-43-3, Polyglutamine
RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence);
BPR (Biological process); BSU (Biological study, unclassified); BIOL
(Biological study); OCCU (Occurrence); PROC (Process)
 (involvement of polyglutamine aggregation in CAG repeat diseases and neuronal cell death)

RN 26700-71-0 HCAPLUS

CN L-Glutamine, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 56-85-9 CMF C5 H10 N2 O3

Absolute stereochemistry.

RN 69864-43-3 HCAPLUS

CN Poly[imino[(1S)-1-(3-amino-3-oxopropyl)-2-oxo-1,2-ethanediyl]] (9CI) (CA INDEX NAME)

IT 101985-79-9, DCAG

RL: ADV (Adverse effect, including toxicity); BIOL (Biological study) (repeat; involvement of polyglutamine aggregation in CAG repeat diseases and neuronal cell death)

RN 101985-79-9 HCAPLUS

CN Guanosine, 2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

L71 ANSWER 6 OF 11 HCAPLUS COPYRIGHT 2003 ACS

AN 1998:272230 HCAPLUS

DN 129:53015

ΤI Tissue transglutaminase-catalyzed formation of high-molecular-weight aggregates in vitro is favored with long polyglutamine domains: a possible mechanism contributing to CAG-triplet diseases

Gentile, Vittorio; Sepe, Carlo; Calvani, Menotti; Melone, Mariarosa A. B.; ΑU Cotrufo, Roberto; Cooper, Arthur J. L.; Blass, John P.; Peluso, Gianfranco

Dipartimento di Biochimica e Biofisica, Seconda Universita di Napoli, CS

Naples, 80138, Italy

Archives of Biochemistry and Biophysics (1998), 352(2), 314-321 SO

CODEN: ABBIA4; ISSN: 0003-9861

PB Academic Press

DT Journal LA English

CC 14-10 (Mammalian Pathological Biochemistry)

To investigate possible biochem. mechanisms underlying the "toxic gain of AΒ function" assocd. with polyglutamine expansions, the ability of guinea pig liver tissue transglutaminase to catalyze covalent attachments of various polyamines to polyglutamine peptides was examd. Of the polyamines tested, spermine is the most active substrate, followed by spermidine and putrescine. Formation of covalent crosslinks between polyglutamine peptides and polyamines yields high-Mr aggregates - a process that is favored with longer polyglutamines. In the presence of tissue transglutaminase, purified glyceraldehyde-3-phosphate dehydrogenase (a key glycolytic enzyme that binds tightly to the polyglutamine domains of both huntingtin and dentatorubralpallidoluysian atrophy proteins) is covalently attached to polyglutamine peptides in vitro, resulting in the formation of high-Mr $\,$ aggregates. In addn., endogenous glyceraldehyde-3-phosphate dehydrogenase of a Balb-c 3T3 fibroblast cell line overexpressing human tissue transglutaminase forms crosslinks with a Q60 polypeptide added to the cell homogenate. Possibly, expansion of polyglutamine domains (thus far known to occur in the gene products assocd. with at least seven neurodegenerative diseases) leads to increased/aberrant tissue transglutaminase-catalyzed crosslinking reactions with both polyamines and susceptible proteins, such as glyceraldehyde-3-phosphate dehydrogenase. Formation of crosslinked heteropolymers may lead to deposition of high-Mr protein aggregates, thereby contributing to cell

ST polyglutamine protein crosslinking aggregation CAG disease; CAG triplet disease polyglutamine protein crosslinking

IT Crosslinking

(biol.; tissue transglutaminase-catalyzed formation of high-mol.-wt. aggregates in vitro is favored with long polyglutamine domains: possible mechanism contributing to human CAG-triplet neurodegenerative diseases)

IT Nervous system

(degeneration; tissue transglutaminase-catalyzed formation of high-mol.-wt. aggregates in vitro is favored with long polyglutamine domains: possible mechanism contributing to human CAG-triplet neurodegenerative diseases)

IT Mutation

(expansion, of CAG trinucleotide repeat; tissue transglutaminase-catalyzed formation of high-mol.-wt. aggregates in vitro is favored with long polyglutamine domains: possible mechanism contributing to human CAG-triplet neurodegenerative diseases)

IT Disease, animal

(genetic, trinucleotide repeat; tissue transglutaminase -catalyzed formation of high-mol.-wt. aggregates in vitro is favored with long polyglutamine domains: possible mechanism contributing to human CAG-triplet neurodegenerative diseases)

IT Amines, biological studies

RL: ADV (Adverse effect, including toxicity); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(polyamines, nonpolymeric, crosslinking; tissue transglutaminase-catalyzed formation of high-mol.-wt. aggregates in vitro is favored with long polyglutamine domains: possible mechanism contributing to human CAG-triplet neurodegenerative diseases)

IT Proteins, specific or class
RL: ADV (Adverse effect, including toxicity); BPR (Biological process);

ΙT

IT

IT

IT

TΤ

ΙT

ΙT

```
BSU (Biological study, unclassified); BIOL (Biological study); PROC
     (Process)
        (polyglutamine-contq.; tissue transglutaminase-catalyzed
        formation of high-mol.-wt. aggregates in vitro is favored with long
       polyglutamine domains: possible mechanism contributing to human
       CAG-triplet neurodegenerative diseases)
    Repeat motifs (protein)
        (polyglutamine; tissue transglutaminase-catalyzed formation
        of high-mol.-wt. aggregates in vitro is favored with long polyglutamine
        domains: possible mechanism contributing to human CAG-triplet
       neurodegenerative diseases)
    Cell death
    Molecular association
        (tissue transglutaminase-catalyzed formation of high-mol.-wt.
        aggregates in vitro is favored with long polyglutamine domains:
       possible mechanism contributing to human CAG-triplet neurodegenerative
       diseases)
    Repetitive DNA
    RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence);
    BSU (Biological study, unclassified); BIOL (Biological study); OCCU
     (Occurrence)
        (trinucleotide; tissue transglutaminase-catalyzed formation
       of high-mol.-wt. aggregates in vitro is favored with long polyglutamine
       domains: possible mechanism contributing to human CAG-triplet
       neurodegenerative diseases)
                                              124-20-9, Spermidine
    71-44-3, Spermine 110-60-1, Putrescine
    9001-50-7, Glyceraldehyde-3-phosphate dehydrogenase
    RL: ADV (Adverse effect, including toxicity); BPR (Biological process);
    BSU (Biological study, unclassified); BIOL (Biological study); PROC
     (Process)
        (crosslinking; tissue transglutaminase-catalyzed formation of
       high-mol.-wt. aggregates in vitro is favored with long polyglutamine
       domains: possible mechanism contributing to human CAG-triplet
       neurodegenerative diseases)
    80146-85-6, Glutaminylpeptide .gamma.-glutamyltransferase
    RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or
    effector, except adverse); BSU (Biological study, unclassified); BIOL
     (Biological study)
        (tissue transglutaminase-catalyzed formation of high-mol.-wt.
        aggregates in vitro is favored with long polyglutamine domains:
       possible mechanism contributing to human CAG-triplet neurodegenerative
       diseases)
    26700-71-0D, Polyglutamine, proteins contg. 69864-43-3D,
    Polyglutamine, proteins contg.
    RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence);
    BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); OCCU (Occurrence); PROC (Process)
        (tissue transglutaminase-catalyzed formation of high-mol.-wt.
        aggregates in vitro is favored with long polyglutamine domains:
       possible mechanism contributing to human CAG-triplet neurodegenerative
       diseases)
    101985-79-9, d-CAG
    RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence);
    BSU (Biological study, unclassified); BIOL (Biological study); OCCU
     (Occurrence)
        (tissue transglutaminase-catalyzed formation of high-mol.-wt.
        aggregates in vitro is favored with long polyglutamine domains:
        possible mechanism contributing to human CAG-triplet neurodegenerative
        diseases)
              THERE ARE 53 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
       53
(1) Aeschlimann, D; Thromb Haemostasis 1994, V71, P402 HCAPLUS
```

(2) Albin, R; Trends Neurosci 1995, V18, P11 HCAPLUS

- (3) Arends, M; Int Rev Exp Pathol 1991, V32, P223 MEDLINE
- (4) Ballestar, E; J Biol Chem 1996, V271, P18817 HCAPLUS
- (5) Beal, M; Ann Neurol 1992, V31, P119 HCAPLUS
- (6) Bessert, D; Mol Brain Res 1995, V33, P165 HCAPLUS
- (7) Browne, S; Ann Neurol 1997, V41, P646 HCAPLUS
- (8) Burke, J; Nature Med 1996, V2, P347 HCAPLUS
- (9) Cooper, A; J Neurochem 1997, V69, P431 HCAPLUS
- (10) Cooper, A; Proc Natl Acad Sci USA 1997, V94, P12604 HCAPLUS
- (11) Davies, S; Cell 1997, V90, P537 HCAPLUS
- (12) Difiglia, M; Neuron 1995, V14, P1075 HCAPLUS
- (13) Esposito, C; J Neurochem 1995, V65, P420 HCAPLUS
- (14) Fesus, L; FEBS Lett 1989, V245, P150 HCAPLUS
- (15) Folk, J; Adv Enzymol 1983, V54, P1 HCAPLUS
- (16) Gentile, V; J Cell Biol 1991, V119, P463
- (17) Green, H; Cell 1993, V74, P955 HCAPLUS
- (18) Greenberg, C; FASEB J 1991, V5, P3071 HCAPLUS
- (19) Grootjans, J; J Biol Chem 1995, V270, P22855 HCAPLUS
- (20) Gutekuns, C; Proc Natl Acad Sci USA 1995, V92, P8710
- (21) Hohenadl, C; J Biol Chem 1995, V270, P23415 HCAPLUS
- (22) Housman, D; Nature Genet 1995, V10, P3 HCAPLUS
- (23) Ikeda, H; Nature Genet 1996, V13, P196 HCAPLUS
- (24) Ishitani, R; J Neurochem 1996, V66, P928 HCAPLUS
- (25) Jenkins, B; Neurology 1993, V43, P2689 MEDLINE
- (26) Jou, Y; Hum Mol Genet 1995, V4, P465 HCAPLUS
- (27) Kahlem, P; Proc Natl Acad Sci USA 1996, V93, P14580 HCAPLUS
- (28) Koshy, B; Hum Mol Genet 1996, V5, P1311 HCAPLUS
- (29) Laemmli, U; Nature 1970, V277, P680
- (30) Lescure, A; EMBO J 1994, V13, P1166 HCAPLUS
- (31) Li, X; Proc Natl Acad Sci USA 1996, V93, P4839 HCAPLUS
- (32) Lorand, L; Ann N Y Acad Sci 1972, V202, P6 HCAPLUS
- (33) Mandel, J; Nature Genet 1994, V7, P453 HCAPLUS
- (34) Mangiarini, L; Cell 1996, V87, P493 HCAPLUS
- (35) Melino, G; Mol Cell Biol 1994, V14, P6584 HCAPLUS
- (36) Penney, J; Ann Neurol 1997, V41, P689
- (37) Perutz, M; Proc Natl Acad Sci USA 1994, V91, P5355 HCAPLUS
- (38) Piacentini, M; Apoptosis II: The Molecular Basis of Apoptosis in Disease 1994, P143 HCAPLUS
- (39) Porta, R; Anal Biochem 1988, V172, P499 HCAPLUS
- (40) Porta, R; Neuropeptides 1988, V11, P89 HCAPLUS
- (41) Porta, R; Phytochemistry 1990, V29, P2801 HCAPLUS
- (42) Portera-Cailliau, C; J Neurosci 1995, V15, P3775 HCAPLUS
- (43) Roses, A; Nature Med 1996, V2, P267 HCAPLUS
- (44) Sambrook, J; Molecular Cloning: A Laboratory Manual 1989
- (45) Saunders, P; J Neurochem 1997, V69, P1820 HCAPLUS
- (46) Sawa, A; Proc Natl Acad Sci USA 1997, V94, P11669 HCAPLUS
- (47) Scherzinger, E; Cell 1997, V90, P549 HCAPLUS
- (48) Singh, U; Biochemistry 1995, V34, P15863 HCAPLUS
- (49) Stott, K; Proc Natl Acad Sci USA 1995, V92, P6509 HCAPLUS
- (50) The Huntington's Disease Collaborative Research Group; Cell 1993, V72, P971
- (51) Thomas, L; Exp Neurol 1995, V133, P265 MEDLINE
- (52) Trottier, Y; Nature Genet 1995, V10, P104 HCAPLUS
- (53) Zeitlin, S; Nature Genet 1995, V11, P155 HCAPLUS
- IT 110-60-1, Putrescine
 - RL: ADV (Adverse effect, including toxicity); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)
 - (crosslinking; tissue **transglutaminase**-catalyzed formation of high-mol.-wt. aggregates in vitro is favored with long polyglutamine domains: possible mechanism contributing to human CAG-triplet neurodegenerative diseases)
- RN 110-60-1 HCAPLUS
- CN 1,4-Butanediamine (8CI, 9CI) (CA INDEX NAME)

 $H_2N-(CH_2)_4-NH_2$

B0146-85-6, Glutaminylpeptide .gamma.-glutamyltransferase RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(tissue transglutaminase-catalyzed formation of high-mol.-wt. aggregates in vitro is favored with long polyglutamine domains: possible mechanism contributing to human CAG-triplet neurodegenerative diseases)

RN 80146-85-6 HCAPLUS

CN Glutamyltransferase, glutaminylpeptide .gamma.- (9CI) (CA INDEX NAME)

26700-71-0D, Polyglutamine, proteins contg. 69864-43-3D,

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

Polyglutamine, proteins contg.
RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence);
BPR (Biological process); BSU (Biological study, unclassified); BIOL

(Biological study); OCCU (Occurrence); PROC (Process)
 (tissue transglutaminase-catalyzed formation of high-mol.-wt.
 aggregates in vitro is favored with long polyglutamine domains:
 possible mechanism contributing to human CAG-triplet neurodegenerative diseases)

RN 26700-71-0 HCAPLUS

CN L-Glutamine, homopolymer (9CI) (CA INDEX NAME)

CM 1

ΙT

CRN 56-85-9. CMF C5 H10 N2 O3

Absolute stereochemistry.

RN 69864-43-3 HCAPLUS

CN Poly[imino[(1S)-1-(3-amino-3-oxopropyl)-2-oxo-1,2-ethanediyl]] (9CI) (CFINDEX NAME)

IT 101985-79-9, d-CAG

RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU

(Occurrence)

(tissue transglutaminase-catalyzed formation of high-mol.-wt. aggregates in vitro is favored with long polyglutamine domains: possible mechanism contributing to human CAG-triplet neurodegenerative diseases)

RN 101985-79-9 HCAPLUS

CN Guanosine, 2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

```
129:3545
DN
     Transglutaminase action imitates Huntington's disease:
ΤI
     selective polymerization of huntingtin containing expanded polyglutamine
     Kahlem, Pascal; Green, Howard; Djian, Philippe
ΑU
     Centre National de la Recherche Scientifique, Centre de Recherche sur
CS
     l'Endocrinologie Moleculaire et le Developpement, Meudon-Bellevue, 92190,
     Molecular Cell (1998), 1(4), 595-601
SO
     CODEN: MOCEFL; ISSN: 1097-2765
PB
     Cell Press
DT
     Journal
LA
     English
     14-10 (Mammalian Pathological Biochemistry)
CC
     Different proteins bearing polyglutamine of excessive length are lethal to
AΒ
     neurons and cause human disease of the central nervous system. In parts
     of the brain affected by Huntington's disease, the amt. of the
     huntingtin with expanded polyglutamine is reduced and there appear
     huntingtin-contg. polymers of larger mol. wt. We show here that
     huntingtin is a substrate of transglutaminase in vitro and that
     the rate const. of the reaction increases with length of the polyglutamine
     over a range of an order of magnitude. As a result, huntingtin with
     expanded polyglutamine is preferentially incorporated into polymers.
                                                                            Both
     disappearance of the huntingtin with expanded polyglutamine and its
     replacement by polymeric forms are prevented by inhibitors of
     transglutaminase. The effect of transglutaminase
     therefore duplicates the changes in the affected parts of the brain.
     Huntington disease huntingtin polyglutamine
ST
     transglutaminase
ΙT
     Nervous system
        (Huntington's chorea; transglutaminase)
        action imitates Huntington's disease by inducing selective
        polymn. of huntingtin contg. expanded polyglutamine)
     Proteins, specific or class
IT
     RL: ADV (Adverse effect, including toxicity); BPR (Biological process);
     BSU (Biological study, unclassified); BIOL (Biological study); PROC
     (Process)
        (huntingtins; transglutaminase action imitates
        Huntington's disease by inducing selective polymn. of
        huntingtin contg. expanded polyglutamine)
     Disease models
TΤ
        (transglutaminase action imitates Huntington's
        disease by inducing selective polymn. of huntingtin contg. expanded
        polyglutamine)
IT
     80146-85-6, Transglutaminase
     RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or
     effector, except adverse); BSU (Biological study, unclassified); BIOL
     (Biological study)
        (transglutaminase action imitates Huntington's
        disease by inducing selective polymn. of huntingtin contg. expanded
        polyglutamine)
     26700-71-0, Polyglutamine 69864-43-3, Polyglutamine
     RL: ADV (Adverse effect, including toxicity); BPR (Biological process);
     BSU (Biological study, unclassified); BIOL (Biological study); PROC
     (Process)
        (transglutaminase action imitates Huntington's
        disease by inducing selective polymn. of huntingtin contg. expanded
        polyglutamine)
     51-85-4, Cystamine
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (transglutaminase action imitates Huntington's
        disease by inducing selective polymn. of huntingtin contg. expanded
```

polyglutamine)

```
THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 35
RE
(1) Aeschlimann, D; Thromb Haemost 1994, V71, P402 HCAPLUS
(2) Aronin, N; Neuron 1995, V15, P1193 HCAPLUS
(3) Cooper, A; J Neurochem 1997, V69, P431 HCAPLUS
(4) Davies, S; Cell 1997, V90, P537 HCAPLUS
(5) de Rooij, K; Human Genet 1995, V95, P270 MEDLINE
(6) Difiglia, M; Science 1997, V277, P1990 HCAPLUS
(7) Goldberg, Y; Nat Genet 1996, V13, P442 HCAPLUS
(8) Green, H; Cell 1993, V74, P955 HCAPLUS
(9) Green, H; In press 1998
(10) Housman, D; Nat Genet 1995, V10, P3 HCAPLUS
(11) Huntington's Disease Collaborative Research Group; Cell 1993, V72, P971
    HCAPLUS
(12) Ikeda, H; Nat Genet 1996, V13, P196 HCAPLUS
(13) Kahlem, P; Proc Natl Acad Sci USA 1996, V93, P14580 HCAPLUS
(14) Laemmli, U; Nature 1970, V227, P680 HCAPLUS
(15) Lorand, L; Proc Natl Acad Sci USA 1996, V93, P14310 HCAPLUS
(16) Mangiarini, L; Cell 1996, V87, P493 HCAPLUS
(17) Ohashi, H; J Biochem 1995, V118, P1271 HCAPLUS
(18) Paulson, H; Neuron 1997, V19, P333 HCAPLUS
(19) Persichetti, F; Mol Med 1995, V1, P374 HCAPLUS
(20) Perutz, M; Proc Natl Acad Sci USA 1994, V91, P5355 HCAPLUS
(21) Reichelt, K; J Neurochem 1992, V59, P500 HCAPLUS
(22) Ross, C; Neuron 1995, V15, P493 HCAPLUS
(23) Scherzinger, E; Cell 1997, V90, P549 HCAPLUS
(24) Schilling, G; Hum Mol Genet 1995, V4, P1365 HCAPLUS
(25) Servadio, A; Nat Genet 1995, V10, P94 HCAPLUS
(26) Sharp, A; Neurobiol Dis 1996, V3, P3 HCAPLUS
(27) Siefring, G; Biochemistry 1978, V17, P2598 HCAPLUS
(28) Simon, M; J Biol Chem 1988, V263, P18093 HCAPLUS
(29) Skinner, P; Nature 1997, V389, P971 HCAPLUS
(30) Stott, K; Proc Natl Acad Sci USA 1995, V92, P6509 HCAPLUS
(31) Telenius, H; Nat Genet 1994, V6, P409 HCAPLUS
(32) Tellez-Nagel, I; J Neuropath Exp Neurol 1974, V33, P308 MEDLINE
(33) Trottier, Y; Nat Genet 1995, V10, P104 HCAPLUS
(34) Vonsattel, J; J Neuropath Exp Neurol 1985, V44, P559 MEDLINE
(35) White, J; Nat Genet 1997, V17, P404 HCAPLUS
     80146-85-6, Transglutaminase
     RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or
     effector, except adverse); BSU (Biological study, unclassified); BIOL
     (Biological study)
        (transglutaminase action imitates Huntington's
        disease by inducing selective polymn. of huntingtin contg. expanded
        polyglutamine)
RN
     80146-85-6 HCAPLUS
     Glutamyltransferase, glutaminylpeptide .gamma. - (9CI) (CA INDEX NAME)
CN
    STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     26700-71-0, Polyglutamine 69864-43-3, Polyglutamine
ΙT
     RL: ADV (Adverse effect, including toxicity); BPR (Biological process);
     BSU (Biological study, unclassified); BIOL (Biological study); PROC
     (Process)
         (transglutaminase action imitates Huntington's
        disease by inducing selective polymn. of huntingtin contg. expanded
        polyglutamine)
RN
     26700-71-0 HCAPLUS
     L-Glutamine, homopolymer (9CI) (CA INDEX NAME)
CN
     CM
          1
     CRN
          56-85-9
```

CMF

C5 H10 N2 O3

Absolute stereochemistry.

RN 69864-43-3 HCAPLUS

CN Poly[imino[(1S)-1-(3-amino-3-oxopropyl)-2-oxo-1,2-ethanediyl]] (9CI) (CA INDEX NAME)

IT **51-85-4**, Cystamine

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(transglutaminase action imitates Huntington's

disease by inducing selective polymn. of huntingtin contg. expanded polyglutamine)

RN 51-85-4 HCAPLUS

CN Ethanamine, 2,2'-dithiobis- (9CI) (CA INDEX NAME)

L71 ANSWER 8 OF 11 HCAPLUS COPYRIGHT 2003 ACS

AN 1998:85454 HCAPLUS

DN 128:179041

TI Suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with an expanded polyglutamine stretch

AU Igarashi, Shuichi; Koide, Reiji; Shimohata, Takayoshi; Yamada, Mitsunori; Hayashi, Yasuko; Takano, Hiroki; Date, Hidetoshi; Oyake, Mutsuo; Sato, Toshiya; Sato, Aki; Egawa, Shigekimi; Ikeuchi, Takeshi; Tanaka, Hajime; Nakano, Ryoichi; Tanaka, Keiko; Hozumi, Isao; Inuzuka, Takashi; Takahashi, Hitoshi; Tsuji, Shoji

CS Dep. Neurology, Niigata Univ., Niigata, 1-757, Japan

SO Nature Genetics (1998), 18(2), 111-117 CODEN: NGENEC; ISSN: 1061-4036

PB Nature America

DT Journal

LA English

CC 14-14 (Mammalian Pathological Biochemistry) Section cross-reference(s): 1, 3

AB To elucidate the mol. mechanisms whereby expanded polyglutamine stretches elicit a grain of toxic function, we expressed full-length and truncated DRPLA (dentatorubral-pallidoluysian atrophy)

cDNAs with or without expanded CAG repeats in COS-7 cells. We found that truncated DRPLA proteins contg. an expanded polyglutamine stretch form filamentous peri- and intranuclear aggregates and undergo apoptosis. The apoptotic cell death was partially suppressed by the transglutaminase inhibitors cystamine and monodansyl cadaverine (but not putrescine), suggesting involvement of a transglutaminase reaction and providing a potential basis for the development of therapeutic measures for CAG-repeat expansion diseases. DRPLA protein transglutaminase inhibitor apoptosis; CAG repeat DRPLA protein cytotoxicity; dentatorubral pallidoluysian

atrophy
IT Proteins, specific or class

RL: ADV (Adverse effect, including toxicity); BIOL (Biological study) (DRPLA (dentatorubral-pallidoluysian atrophy); suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with expanded polyglutamine stretch)

IT Brain, disease

(dentatorubral-pallidoluysian atrophy; suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with expanded polyglutamine stretch)

IT Apoptosis

ST

(suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with expanded polyglutamine stretch)

IT Repetitive DNA

RL: ADV (Adverse effect, including toxicity); BIOL (Biological study) (trinucleotide; suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with expanded polyglutamine stretch)

IT 80146-85-6, Transglutaminase

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (inhibitors; suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with expanded polyglutamine stretch)

IT **101985-79-9**, d-CAG

RL: ADV (Adverse effect, including toxicity); BIOL (Biological study) (suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with expanded polyglutamine stretch)

IT 51-85-4, Cystamine 10121-91-2, Monodansyl cadaverine

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with expanded polyglutamine stretch)

RE.CNT 38 THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD RE

- (1) Burright, E; Cell 1995, V82, P937 HCAPLUS
- (2) David, G; Nature Genet 1997, V17, P65 HCAPLUS
- (3) Davies, S; Cell 1997, V90, P537 HCAPLUS
- (4) Dickson, R; J Biol Chem 1981, V256, P3454 HCAPLUS
- (5) Difiglia, M; Science 1997, V277, P1990 HCAPLUS
- (6) Goldberg, Y; Nature Genet 1996, V13, P442 HCAPLUS
- (7) Ikeda, H; Nature Genet 1996, V13, P196 HCAPLUS
- (8) Ikeuchi, T; Ann Neurol 1995, V37, P769 MEDLINE
- (9) Imbert, G; Nature Genet 1996, V14, P285 HCAPLUS
- (10) Jackson, M: Neuropatho Appl Neurobiol 1995, V21, P18 MEDLINE
- (11) Kahlem, P; Proc Natl Acad Sci USA 1996, V93, P14580 HCAPLUS
- (12) Kawaguchi, Y; Nature Genet 1994, V8, P221 HCAPLUS
- (13) Kleman, J; Biochemistry 1995, V34, P13768 HCAPLUS
- (14) Koide, R; Nature Genet 1994, V6, P9 HCAPLUS

- (15) La Spada, A; Nature 1991, V352, P77 HCAPLUS (16) Lorand, L; Biochemistry 1979, V18, P1756 HCAPLUS (17) Lubahn, D; Science 1988, V240, P327 HCAPLUS (18) Mangiarini, L; Cell 1996, V87, P493 HCAPLUS (19) Mizushima, S; Nucleic Acids Res 1990, V18, P5322 HCAPLUS (20) Mori, Y; Nature 1991, V350, P398 HCAPLUS (21) Nagafuchi, S; Nature Genet 1994, V6, P14 HCAPLUS (22) Nagafuchi, S; Nature Genet 1994, V8, P177 HCAPLUS (23) Naito, H; Neurol 1982, V32, P798 MEDLINE (24) Onodera, O; Am J Hum Genet 1995, V57, P1050 HCAPLUS (25) Onodera, O; Biochem Biophy Res Commun 1997, V238, P599 HCAPLUS (26) Orr, H; Nature Genet 1993, V4, P221 HCAPLUS (27) Paulson, H; Ann Neurol 1997, V41, P453 HCAPLUS (28) Paulson, H; Neuron 1997, V19, P333 HCAPLUS (29) Perutz, M; Proc Natl Acad Sci USA 1994, V91, P5355 HCAPLUS (30) Pulst, S; Nature Genet 1996, V14, P269 HCAPLUS (31) Sanpei, K; Nature Genet 1996, V14, P277 HCAPLUS (32) Scherzinger, E; Cell 1997, V90, P549 HCAPLUS (33) Servadio, A; Nature Genet 1995, V10, P94 HCAPLUS (34) Stott, K; Proc Natl Acad Sci USA 1995, V92, P6509 HCAPLUS (35) The Huntington's Disease Collaborative Research Group; Cell 1993, V72, P971 (36) Trottier, Y; Nature Genet 1995, V10, P104 HCAPLUS (37) Yazawa, I; Nature Genet 1995, V10, P99 HCAPLUS (38) Zhuchenko, O; Nature Genet 1997, V15, P62 HCAPLUS 80146-85-6, Transglutaminase TΤ RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (inhibitors; suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with expanded polyglutamine stretch) RN 80146-85-6 HCAPLUS Glutamyltransferase, glutaminylpeptide .gamma.- (9CI) (CA INDEX NAME) CN
- *** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
- ΙT **101985-79-9**, d-CAG
 - RL: ADV (Adverse effect, including toxicity); BIOL (Biological study) (suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with expanded polyglutamine stretch)
- RN 101985-79-9 HCAPLUS
- Guanosine, 2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-CN (3'.fwdarw.5')-2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

IT 51-85-4, Cystamine 10121-91-2, Monodansyl cadaverine

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with expanded polyglutamine stretch)

RN 51-85-4 HCAPLUS

CN Ethanamine, 2,2'-dithiobis- (9CI) (CA INDEX NAME)

 $H_2N-CH_2-CH_2-S-S-CH_2-CH_2-NH_2$

- RN 10121-91-2 HCAPLUS
- CN 1-Naphthalenesulfonamide, N-(5-aminopentyl)-5-(dimethylamino)- (8CI, 9CI) (CA INDEX NAME)

- L71 ANSWER 9 OF 11 HCAPLUS COPYRIGHT 2003 ACS
- AN 1997:768662 HCAPLUS
- DN 128:46792
- TI Transglutaminase-catalyzed inactivation of glyceraldehyde 3-phosphate dehydrogenase and .alpha.-ketoglutarate dehydrogenase complex by polyglutamine domains of pathological length
- AU Cooper, Arthur J. L.; Sheu, K. -F. Rex; Burke, James R.; Onodera, Osamu; Strittmatter, Warren J.; Roses, Allen D.; Blass, John P.
- CS Department of Biochemistry, Cornell University Medical College, New York, NY, 10021, USA
- SO Proceedings of the National Academy of Sciences of the United States of America (1997), 94(23), 12604-12609
 CODEN: PNASA6; ISSN: 0027-8424
- PB National Academy of Sciences
- DT Journal
- LA English
- CC 14-10 (Mammalian Pathological Biochemistry)
 Section cross-reference(s): 3
- Several adult-onset neurodegenerative diseases are caused by genes with AΒ expanded CAG triplet repeats within their coding regions and extended polyglutamine (Qn) domains within the expressed proteins. Generally, in clin. affected individuals n .gtoreq. 40. Glyceraldehyde 3-phosphate dehydrogenase binds tightly to four Qn disease proteins, but the significance of this interaction is unknown. The authors now report that purified glyceraldehyde 3-phosphate dehydrogenase is inactivated by tissue transglutaminase in the presence of glutathione S-transferase constructs contg. a Qn domain of pathol. length (n = 62 or 81). dehydrogenase is less strongly inhibited by tissue transglutaminase in the presence of constructs contg. shorter Qn domains (n = 0 or 10). Purified .alpha.-ketoglutarate dehydrogenase complex also is inactivated by tissue transglutaminase plus glutathione S-transferase constructs contg. pathol.-length Qn domains (n = 62 or 81). Apparently, tissue transglutaminase-catalyzed covalent linkages involving the larger poly-Q domains may disrupt cerebral energy metab. in CAG/Qn expansion diseases.
- ST GAPDH inhibition polyglutamine tissue **transglutaminase** neurodegeneration
- IT Gene, animal
 - RL: BSU (Biological study, unclassified); BIOL (Biological study)
 (CAG triplet contg.; longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)
- IT Nervous system

(Huntington's chorea; longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase,

but do so in the presence of tissue transglutaminase)

IT Spinal muscular atrophy

(X-linked spinal and bulbar muscular atrophy; longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

IT Nervous system

(degeneration; longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

IT Brain, disease

(dentatorubral-pallidoluysian atrophy; longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

IT Metabolism

(energy; longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

IT Disease, animal

(genetic, expansion diseases; longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

IT Protein motifs

(polyglutamine domain; longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue **transglutaminase**)

IT Nervous system

(spinocerebellar ataxia; longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

IT 9001-50-7, Glyceraldehyde-3-phosphate dehydrogenase **26700-71-0**, Polyglutamine

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study).

(longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

IT 80146-85-6, Tissue transglutaminase

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

IT 101985-79-9

RL: BSU (Biological study, unclassified); BIOL (Biological study) (longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

IT 26700-71-0, Polyglutamine

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

RN 26700-71-0 HCAPLUS

CN L-Glutamine, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 56-85-9 CMF C5 H10 N2 O3 Absolute stereochemistry.

IT 80146-85-6, Tissue transglutaminase

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue **transglutaminase**)

RN 80146-85-6 HCAPLUS

CN Glutamyltransferase, glutaminylpeptide .gamma.- (9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

IT 101985-79-9

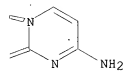
RL: BSU (Biological study, unclassified); BIOL (Biological study) (longer polyglutamine domains by themselves do not inhibit glyceraldehyde 3-phosphate dehydrogenase, but do so in the presence of tissue transglutaminase)

RN 101985-79-9 HCAPLUS

CN Guanosine, 2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxy-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B



L71 ANSWER 10 OF 11 HCAPLUS COPYRIGHT 2003 ACS

AN 1997:409539 HCAPLUS

DN 127:134212

TI Polyglutamine domains are substrates of tissue transglutaminase: does transglutaminase play a role in expanded CAG/Poly-Q neurodegenerative diseases?

AU Cooper, Arthur J. L.; Sheu, Kwan-Fu Rex; Burke, James R.; Onodera, Osamu; Strittmatter, Warren J.; Roses, Allen D.; Blass, John P.

CS Department of Biochemistry, Cornell University Medical College, New York, NY, USA

SO Journal of Neurochemistry (1997), 69(1), 431-434 CODEN: JONRA9; ISSN: 0022-3042

PB Lippincott-Raven

DT Journal

LA English

CC 14-10 (Mammalian Pathological Biochemistry) Section cross-reference(s): 7

Huntington's disease and six other neurodegenerative diseases are assocd. with abnormal gene products contg. expanded polyglutamine (poly-Q; Qn) domains (n .gtoreq. 40). In the present work, the authors show that glutathione S-transferase (GST) fusion proteins contg. a small, physiol.-length poly-Q domain (GSTQ10) or a large, pathol.-length poly-Q domain (GSTQ62) are excellent substrates of guinea pig liver (tissue) transglutaminase and that both GSTQ10 and GSTQ62 are activators of tissue transglutaminase-catalyzed hydroxaminolysis of N-.alpha.-carbobenzoxyglutaminylglycine. The present findings have implications for understanding the pathophysiol. mechanisms of expanded CAG/poly-Q domain diseases.

ST transglutaminase polyglutamine protein substrate neurodegenerative disease; Huntington transglutaminase polyglutamine protein substrate

IT Gene, animal

RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence)

(CAG repeat-contg.; proteins contg. polyglutamine domains are substrates of tissue **transglutaminase** and **transglutaminase** may play role in human expanded

CAG/polyglutamine neurodegenerative diseases) ΙT Nervous system (Huntington's chorea; proteins contg. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) ΙT Nervous system (degeneration, trinucleotide repeat; proteins contq. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) ΙT Proteins, specific or class RL: ADV (Adverse effect, including toxicity); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (expanded polyglutamine-contg.; proteins contg. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) ΙT Mutation (expansion, of CAG trinucleotide repeat; proteins contq. polyqlutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) ΙT Repeat motifs (protein) RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence); PROC (Process) (polyglutamine; proteins contg. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) ΙT Repetitive DNA RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence) (trinucleotide; proteins contg. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) 80146-85-6, Glutaminylpeptide .gamma.-glutamyltransferase ΙT RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (proteins contg. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) 26700-71-0D, Polyglutamine, proteins contg. 69864-43-3D, ITPolyglutamine, proteins contg. RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence); PROC (Process) (proteins contg. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) ΙT 101985-79-9, d-CAG RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence) (proteins contg. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) 80146-85-6, Glutaminylpeptide .gamma.-glutamyltransferase ΙT

RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (proteins contg. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) RN . 80146-85-6 HCAPLUS Glutamyltransferase, glutaminylpeptide .gamma.- (9CI) (CA INDEX NAME) CN*** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 26700-71-0D, Polyglutamine, proteins contg. 69864-43-3D, Polyglutamine, proteins contg. RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence); PROC (Process) (proteins contg. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role in human expanded CAG/polyglutamine neurodegenerative diseases) RN 26700-71-0 HCAPLUS L-Glutamine, homopolymer (9CI) (CA_INDEX NAME) CN

CM 1

CRN 56-85-9 CMF C5 H10 N2 O3

Absolute stereochemistry.

RN 69864-43-3 HCAPLUS
CN Poly[imino[(1S)-1-(3-amino-3-oxopropyl)-2-oxo-1,2-ethanediyl]] (9CI) ((INDEX NAME)

IT **101985-79-9**, d-CAG

RL: ADV (Adverse effect, including toxicity); BOC (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence)

(proteins contg. polyglutamine domains are substrates of tissue transglutaminase and transglutaminase may play role

in human expanded CAG/polyglutamine neurodegenerative diseases) RN 101985-79-9 HCAPLUS

CN Guanosine, 2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxy-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

L71 ANSWER 11 OF 11 HCAPLUS COPYRIGHT 2003 ACS

AN ' 1996:746875 HCAPLUS

DN 126:101006

ΤI Peptides containing glutamine repeats as substrates for transglutaminase-catalyzed crosslinking: relevance to diseases of the nervous system

ΑU

Kahlem, P.; Terre, C.; Green, H.; Djian, P. Cent. Natl. Rech. Sci., Cent. Rech. Endocrinol. Mol. Dev., CS Meudon-Bellevue, 92190, Fr.

Proceedings of the National Academy of Sciences of the United States of SO America (1996), 93(25), 14580-14585

2

CODEN: PNASA6; ISSN: 0027-8424 PΒ National Academy of Sciences DTJournal LA English CC 7-3 (Enzymes) Many proteins contain reiterated glutamine residues, but polyglutamine of AB excessive length may result in human disease by conferring new properties on the protein contq. it. One established property of a glutamine residue, depending on the nature of the flanking residues, is its ability to act as an amine acceptor in a transglutaminase-catalyzed reaction and to make a glutamyl-lysine cross-link with a neighboring To learn whether glutamine repeats can act as amine polypeptide. acceptors, we have made peptides with variable lengths of polyglutamine flanked by the adjacent amino acid residues in the proteins assocd. with spinocerebellar ataxia type 1 (SCA1), Machado-Joseph disease (SCA3), or dentato-rubral pallidoluysian atrophy (DRPLA) or those residues adjacent to the preferred crosslinking site of involucrin, or solely by arginine residues. The polyglutamine was found to confer excellent substrate properties on any sol. peptide; under optimal conditions, virtually all the glutamine residues acted as amine acceptors in the reaction with glycine ethyl-ester, and lengthening the sequence of polyglutamine increased the reactivity of each glutamine residue. In the presence of transglutaminase, peptides contg. polyglutamine formed insol. aggregates with the proteins of brain exts. and these aggregates contained glutamyl-lysine cross-links. Repeated glutamine residues exposed on the surface of a neuronal protein should form cross-linked aggregates in the presence of any transglutaminase activated by the presence of Ca2+. transglutaminase substrate glutamine calcium STITBrain, disease RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); PROC (Process) (dentatorubral-pallidoluysian atrophy; peptides contq. glutamine repeats as substrates for transglutaminase -catalyzed crosslinking and relevance to nervous system diseases) IT Nervous system RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); PROC (Process) (disease, spinocerebellar ataxia 3; peptides contg. glutamine repeats as substrates for transglutaminase -catalyzed crosslinking and relevance to nervous system diseases) ΙT Proteins, specific or class RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); PROC (Process) (involucrins; peptides contg. glutamine repeats as substrates for transglutaminase-catalyzed crosslinking and relevance to nervous system diseases) ΙT Nervous system RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); PROC (Process) (spinocerebellar ataxia 1; peptides contg. glutamine repeats as substrates for transglutaminase -catalyzed crosslinking and relevance to nervous system diseases) ΙT Structure-activity relationship (transglutaminase substrates; peptides contg. glutamine repeats as substrates for transglutaminase-catalyzed crosslinking) ΙT 7440-70-2, Calcium, biological studies 80146-85-6, Transglutaminase RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study) (peptides contg. glutamine repeats as substrates for

transglutaminase-catalyzed crosslinking)

56-85-9, Glutamine, biological studies RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (peptides contg. glutamine repeats as substrates for transglutaminase-catalyzed crosslinking) 26700-71-0, Polyglutamine 69864-43-3, Polyglutamine IT RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); PROC (Process) (peptides contg. glutamine repeats as substrates for transglutaminase-catalyzed crosslinking) ΙT 80146-85-6, Transglutaminase RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study) (peptides contg. glutamine repeats as substrates for transglutaminase-catalyzed crosslinking) RN 80146-85-6 HCAPLUS Glutamyltransferase, glutaminylpeptide .gamma.- (9CI) (CA INDEX NAME) CN *** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 26700-71-0, Polyglutamine 69864-43-3, Polyglutamine IT RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); PROC (Process) (peptides contg. glutamine repeats as substrates for transglutaminase-catalyzed crosslinking) RN 26700-71-0 HCAPLUS CN L-Glutamine, homopolymer (9CI) (CA INDEX NAME) CM1 CRN 56-85-9

Absolute stereochemistry.

CMF

C5 H10 N2 O3

RN 69864-43-3 HCAPLUS
CN Poly[imino[(1S)-1-(3-amino-3-oxopropyl)-2-oxo-1,2-ethanediyl]] (9CI) (CAINDEX NAME)

=> sel hit rn E1 THROUGH E14 ASSIGNED

: 20

=> fil reg FILE 'REGISTRY' ENTERED AT 16:08:12 ON 01 JUL 2003 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2003 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

30 JUN 2003 HIGHEST RN 540462-79-1 STRUCTURE FILE UPDATES: 30 JUN 2003 HIGHEST RN 540462-79-1 DICTIONARY FILE UPDATES:

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

=> s e1-e14

(80146-85-6/RN) 1 26700-71-0/BI (26700-71-0/RN) 1 69864-43-3/BI (69864-43-3/RN) 1 101985-79-9/BI (101985-79-9/RN) 1 51-85-4/BI (51-85-4/RN)1 10121-91-2/BI (10121-91-2/RN) 1 110-60-1/BI (110-60-1/RN)1 150-13-0/BI (150-13-0/RN)1 24991-23-9/BI (24991-23-9/RN) 1 25513-46-6/BI (25513-46-6/RN)1 616-34-2/BI (616-34-2/RN)1 64-77-7/BI (64-77-7/RN)1 74389-76-7/BI (74389-76-7/RN) 1 7758-98-7/BI (7758-98-7/RN)

1 80146-85-6/BI

14 (80146-85-6/BI OR 26700-71-0/BI OR 69864-43-3/BI OR 101985-79-9/ BI OR 51-85-4/BI OR 10121-91-2/BI OR 110-60-1/BI OR 150-13-0/BI OR 24991-23-9/BI OR 25513-46-6/BI OR 616-34-2/BI OR 64-77-7/BI OR 74389-76-7/BI OR 7758-98-7/BI)

=> d ide can tot

L72

ANSWER 1 OF 14 REGISTRY COPYRIGHT 2003 ACS L72 RN 101985-79-9 REGISTRY CN

Guanosine, 2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-

(3'.fwdarw.5')-2'-deoxy- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 18: PN: US20030093830 PAGE: 18 claimed sequence

CN 99: PN: WO02072882 TABLE: 3 claimed sequence

CN DCAG

CN Deoxy-CAG trinucleotide

FS STEREOSEARCH

MF C29 H37 N13 O15 P2

·CI COM

SR CA

LC STN Files: BEILSTEIN*, CA, CAPLUS, TOXCENTER, USPATFULL (*File contains numerically searchable property data)

Absolute stereochemistry.

PAGE 1-B

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT 497 REFERENCES IN FILE CA (1957 TO DATE) 10 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA 499 REFERENCES IN FILE CAPLUS (1957 TO DATE) REFERENCE 1: 139:4926 REFERENCE 139:1618 138:383374 REFERENCE 138:380372 REFERENCE 138:366988 REFERENCE 5: REFERENCE 6: 138:366585 7: 138:366424 REFERENCE 138:366368 REFERENCE 8: 138:319127 REFERENCE 9: 10: 138:301261 REFERENCE L72 ANSWER 2 OF 14 REGISTRY COPYRIGHT 2003 ACS **80146-85-6** REGISTRY RN Glutamyltransferase, glutaminylpeptide .gamma. - (9CI) (CA INDEX NAME) CN OTHER NAMES: Activa MP CN Activa Supercurd CN Activa TG CN Activa TG-K CNCN Activa TG-M Activa TG-S CN Activa TG-TI CN CN Activa WM CN Akuthiba TG-S CN E.C. 2.3.2.13 Glutaminylpeptide .gamma.-glutamyltransferase CN CN Koshikeep CN Polyamine transglutaminase CN PPQ 6117 CN Tissue transglutaminase ${\tt Transglutaminase}$ CN 300711-04-0 DR Unspecified MF CI MAN ADISNEWS, AGRICOLA, ANABSTR, BIOBUSINESS, BIOSIS, BIOTECHNO, LC STN Files: CA, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMLIST, CIN, EMBASE, MSDS-OHS, PIRA, PROMT, RTECS*, TOXCENTER, USPAT2, USPATFULL (*File contains numerically searchable property data) *** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 2789 REFERENCES IN FILE CA (1957 TO DATE) 27 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA 2799 REFERENCES IN FILE CAPLUS (1957 TO DATE)

REFERENCE

1: 139:6037

REFERENCE 2: 139:5395

REFERENCE 3: 139:4647

REFERENCE 4: 139:2821

REFERENCE 5: 139:2758

REFERENCE 6: 139:2711

REFERENCE 7: 138:400779

REFERENCE 8: 138:400693

REFERENCE 9: 138:399776

REFERENCE 10: 138:399774

L72 ANSWER 3 OF 14 REGISTRY COPYRIGHT 2003 ACS

RN **74389-76-7** REGISTRY

CN Norvaline, 5-diazo-4-oxo-N-[(phenylmethoxy)carbonyl]-, 4-nitrophenyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C19 H16 N4 O7

LC STN Files: CA, CAPLUS, TOXCENTER

4 REFERENCES IN FILE CA (1957 TO DATE)

4 REFERENCES IN FILE CAPLUS (1957 TO DATE)

REFERENCE 1: 132:45002

REFERENCE 2: 97:196229

REFERENCE 3: 94:188083

REFERENCE 4: 93:68458

L72 ANSWER 4 OF 14 REGISTRY COPYRIGHT 2003 ACS

RN **69864-43-3** REGISTRY

CN Poly[imino[(1S)-1-(3-amino-3-oxopropyl)-2-oxo-1,2-ethanediyl]] (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Poly[imino[1-(3-amino-3-oxopropyl)-2-oxo-1,2-ethanediyl]], (S)-OTHER NAMES:

CN Poly(glutamine), SRU

CN Poly(L-glutamine), SRU

CN Poly-L-glutamine

CN Polyglutamine

DR 26603-78-1

MF (C5 · H8 N2 O2) n

CI PMS, COM

PCT Polyamide

LC STN Files: ADISNEWS, AGRICOLA, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CAPLUS, CEN, CIN, EMBASE, PIRA, PROMT, TOXCENTER, USPAT2, USPATFULL

RELATED POLYMERS AVAILABLE WITH POLYLINK

205 REFERENCES IN FILE CA (1957 TO DATE)
19 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

206 REFERENCES IN FILE CAPLUS (1957 TO DATE)

REFERENCE 1: 139:4931

REFERENCE 2: 139:4926

REFERENCE 3: 139:1618

REFERENCE 4: 138:396052

REFERENCE 5: 138:390990

REFERENCE 6: 138:348760

REFERENCE 7: 138:335249

REFERENCE 8: 138:318968

REFERENCE 9: 138:314634

REFERENCE 10: 138:202962

L72 ANSWER 5 OF 14 REGISTRY COPYRIGHT 2003 ACS

RN **26700-71-0** REGISTRY

CN L-Glutamine, homopolymer (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Glutamine, L-, peptides (8CI)

OTHER NAMES:

CN Glutamine homopolymer

CN Poly-L-glutamine

CN Polyglutamine

FS STEREOSEARCH

MF (C5 H10 N2 O3)x

CI PMS, COM

PCT Polyamide, Polyamide formed

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAPLUS, CEN, CIN, EMBASE, MEDLINE, PIRA, PROMT, TOXCENTER, USPAT2, USPATFULL

RELATED POLYMERS AVAILABLE WITH POLYLINK

CRN 56-85-9 CMF C5 H10 N2 O3

Absolute stereochemistry.

558 REFERENCES IN FILE CA (1957 TO DATE)
36 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
560 REFERENCES IN FILE CAPLUS (1957 TO DATE)

REFERENCE 1: 139:4931

REFERENCE 2: 139:4926

REFERENCE 3: 139:1618

REFERENCE 4: 138:399369

REFERENCE 5: 138:397691

REFERENCE 6: 138:396052

REFERENCE 7: 138:390990

REFERENCE 8: 138:382993

REFERENCE 9: 138:382992

REFERENCE 10: 138:380166

L72 ANSWER 6 OF 14 REGISTRY COPYRIGHT 2003 ACS

RN 25513-46-6 REGISTRY

CN L-Glutamic acid, homopolymer (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Glutamic acid, L-, peptides (8CI)

OTHER NAMES:

CN .alpha.-L-Glutamic acid polymer

CN .gamma.-L-Polyglutamic acid

CN Glutamic acid homopolymer

CN Glutamic acid polymer

CN L-Glutamic acid polymer

CN PGA

CN Poly(.alpha.-L-glutamic acid)

CN Poly(L-glutamic acid)

CN Poly-L-glutamate

CN Polyglutamic acid

FS STEREOSEARCH

DR 24938-00-9, 115529-71-0, 66415-63-2, 141982-72-1, 84960-48-5, 26717-13-5

MF (C5 H9 N O4) \times

CI PMS, COM

PCT Polyamide, Polyamide formed

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CIN, CSCHEM, DDFU, DIOGENES, DRUGU, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, PIRA, PROMT, RTECS*, TOXCENTER, TULSA, USPAT2, USPATFULL (*File contains numerically searchable property data)

RELATED POLYMERS AVAILABLE WITH POLYLINK

CM 1

CRN 56-86-0 CMF C5 H9 N O4

Absolute stereochemistry.

1890 REFERENCES IN FILE CA (1957 TO DATE)

391 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

1896 REFERENCES IN FILE CAPLUS (1957 TO DATE)

REFERENCE 1: 139:16809

REFERENCE 2: 139:5708

REFERENCE 3: 139:3049

REFERENCE 4: 139:979

REFERENCE 5: 138:411206

REFERENCE 6: 138:406739

REFERENCE 7: 138:398299

REFERENCE 8: 138:397896

REFERENCE 9: 138:390607

REFERENCE 10: 138:373954

L72 ANSWER 7 OF 14 REGISTRY COPYRIGHT 2003 ACS

RN 24991-23-9 REGISTRY

CN Poly[imino[(1S)-1-(2-carboxyethyl)-2-oxo-1,2-ethanediyl]] (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Poly[iminocarbonyl(3-carboxypropylidene)], L- (8CI)

CN Poly[imino[1-(2-carboxyethyl)-2-oxo-1,2-ethanediyl]], (S)-

OTHER NAMES:

CN Glutamic acid homopolymer, SRU

CN L-Glutamic acid homopolymer, SRU

CN Poly(.alpha.-glutamic acid), SRU

CN Poly(.alpha.-L-glutamic acid), SRU

CN Poly(L-.alpha.-glutamyl)

CN Poly(L-glutamic acid), SRU

CN Poly(L-glutamyl)

CN Poly-L-glutamate SRU

CN Polyglutamic acid, SRU

DR 124224-52-8, 37453-50-2, 78678-46-3, 26063-12-7, 26915-14-0

MF (C5 H7 N O3)n

CI PMS, COM

22

PCT Polyamide

LC STN Files: BIOSIS, BIOTECHNO, CA, CAPLUS, CASREACT, EMBASE, IFICDB, IFIPAT, IFIUDB, TOXCENTER, USPAT2, USPATFULL

RELATED POLYMERS AVAILABLE WITH POLYLINK

```
CH2-CH2-CO2H
NH-CH-
```

1430 REFERENCES IN FILE CA (1957 TO DATE) 320 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

1435 REFERENCES IN FILE CAPLUS (1957 TO DATE)

REFERENCE 1: 139:5708

REFERENCE 139:979

138:406739 REFERENCE

REFERENCE 138:398299

REFERENCE 138:397896

REFERENCE 138:390607

REFERENCE 138:373954

REFERENCE 138:369333

REFERENCE 138:369188

REFERENCE 10: 138:354701

L72 ANSWER 8 OF 14 REGISTRY COPYRIGHT 2003 ACS

10121-91-2 REGISTRY

1-Naphthalenesulfonamide, N-(5-aminopentyl)-5-(dimethylamino)- (8CI, 9CI) CN (CA INDEX NAME)

OTHER NAMES:

CN Dansylcadaverine

Monodansylcadaverine CN

Other Sources:

 $N\hbox{-} (5\hbox{-}Amin opentyl) \hbox{-} 5\hbox{-}dimethylamino-1\hbox{-}naphthalene sulfonamide}$ CN

FS

99473-69-5 DR

MF C17 H25 N3 O2 S

STN Files: ADISNEWS, AGRICOLA, BEILSTEIN*, BIOBUSINESS, BIOSIS, LCBIOTECHNO, CA, CANCERLIT, CAPLUS, CASREACT, CHEMCATS, CHEMLIST, CSCHEM, DDFU, DRUGU, EMBASE, MEDLINE, NIOSHTIC, TOXCENTER, USPATFULL, VETU (*File contains numerically searchable property data) EINECS**

(**Enter CHEMLIST File for up-to-date regulatory information)

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

263 REFERENCES IN FILE CA (1957 TO DATE)

12 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

263 REFERENCES IN FILE CAPLUS (1957 TO DATE)

REFERENCE 1: 138:272300

REFERENCE 2: 138:233597

REFERENCE 138:122793

REFERENCE 137:245263 4:

REFERENCE 5: 137:105614

136:337048 REFERENCE

7: 136:66576 REFERENCE

REFERENCE 136:1616

REFERENCE 135:343279

REFERENCE 10: 135:177036

L72 ANSWER 9 OF 14 REGISTRY COPYRIGHT 2003 ACS

7758-98-7 REGISTRY

Sulfuric acid copper(2+) salt (1:1) (8CI, 9CI) (CA INDEX NAME)

OTHER NAMES:

CN Blue Copper

Blue stone CN

CN Blue vitriol

CN Copper monosulfate

Copper sulfate CN

CN Copper sulfate (1:1)

CNCopper sulfate (CuSO4)

CN Copper(2+) sulfate

Copper(2+) sulfate (1:1) CN

CN Copper(II) sulfate

CN Cuivrol

CN Cupric sulfate

CN Cupric sulfate anhydrous

CN Cupric sulphate

CN Delcup

CN Hylinec

CN Incracide 10A CN Incracide E 51

CN MAC 570

```
CN
    .Monocopper sulfate
CN
     Roman vitriol
     Sulfuric acid, copper(2+) salt (1:1)
CN
DR
     139939-69-8
MF
     Cu . H2 O4 S
CI
     COM
                  ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS,
LC
     STN Files:
       BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS,
       CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM*, DIOGENES, DIPPR*, DRUGU,
       EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT, ENCOMPPAT2, GMELIN*, HSDB*,
       IFICDB, IFIPAT, IFIUDB, MEDLINE, MRCK*, MSDS-OHS, NIOSHTIC, PDLCOM*,
       PIRA, PROMT, RTECS*, TOXCENTER, TULSA, USAN, USPAT2, USPATFULL, VETU,
         (*File contains numerically searchable property data)
                      DSL**, EINECS**, TSCA**
         (**Enter CHEMLIST File for up-to-date regulatory information)
CRN
     (7664 - 93 - 9)
    0
HO-S
     — ОН
    0
  Cu(II)
           17916 REFERENCES IN FILE CA (1957 TO DATE)
             229 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
           17932 REFERENCES IN FILE CAPLUS (1957 TO DATE)
                139:16836
REFERENCE
                139:16113
REFERENCE
            2:
REFERENCE
                139:15101
REFERENCE
                139:14719
REFERENCE
                139:13883
REFERENCE
                139:11485
                139:7579
REFERENCE
            7:
REFERENCE
            8:
                139:6038
REFERENCE
                139:5709
REFERENCE
           10:
                139:2372
     ANSWER 10 OF 14 REGISTRY COPYRIGHT 2003 ACS
RN
     616-34-2 REGISTRY
     Glycine, methyl ester (6CI, 8CI, 9CI) (CA INDEX NAME)
OTHER NAMES:
CN
     (Methoxycarbonyl) methylamine
CN
     Glycine O-methyl ester
CN
     Methyl aminoacetate
CN
     Methyl glycinate
```

Methyl glycine

CN

```
3D CONCORD
FS
MF
     C3 H7 N O2
CI
     COM
                  BEILSTEIN*, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAOLD, CAPLUS,
LC
       CASREACT, CHEMCATS, CHEMINFORMRX, CHEMLIST, EMBASE, GMELIN*, HODOC*,
       IFICDB, IFIPAT, IFIUDB, MEDLINE, SPECINFO, TOXCENTER, USPAT7, USPATFULL
         (*File contains numerically searchable property data)
                      EINECS**
     Other Sources:
         (**Enter CHEMLIST File for up-to-date regulatory information)
    0
MeO-C-CH2-NH2
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
            1002 REFERENCES IN FILE CA (1957 TO DATE)
              44 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
            1005 REFERENCES IN FILE CAPLUS (1957 TO DATE)
              27 REFERENCES IN FILE CAOLD (PRIOR TO 1967)
REFERENCE
                139:7371
                139:6418
REFERENCE
            2:
REFERENCE
            3:
                138:385730
                138:385680
REFERENCE
                138:385420
REFERENCE
                138:353826
REFERENCE
REFERENCE
                138:353778
            7:
REFERENCE
                138:303687
REFERENCE
                138:287984
REFERENCE
           10:
                138:260997
    ANSWER 11 OF 14 REGISTRY COPYRIGHT 2003 ACS
     150-13-0 REGISTRY
                                    (CA INDEX NAME)
     Benzoic acid, 4-amino- (9CI)
OTHER CA INDEX NAMES:
     Benzoic acid, p-amino- (8CI)
OTHER NAMES:
CN
     4-Aminobenzoic acid
CN
     4-Carboxyaniline
CN
CN
     Aniline-4-carboxylic acid
CN
     Anti-Chromotrichia factor
CN
     Anticanitic vitamin
CN
     Anticantic vitamin
CN
     Antichromotrichia factor
CN
     Bacterial vitamin H1
CN
     Chromotrichia factor
CN
     Hachemina
```

CN

CN

p-Aminobenzoic acid

p-Carboxyaniline

```
p-Carboxyphenylamine
CN
CN
     PAB.
CN
     PABA
CN
     Pabacyd
CN
     Pabafilm'
CN
     Pabamine
CN
     Paraminol
CN
     Paranate
CN
     Romavit
CN
     Sunbrella
     Trichochromogenic factor
CN
CN
     Vitamin BX
     Vitamin H'
CN
FS
     3D CONCORD
     8014-65-1
DR
MF
     C7 H7 N O2
CI
     COM
                  ADISINSIGHT, ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BEILSTEIN*,
LC
       BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS,
       CASREACT, CBNB, CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM,
       CSNB, DDFU, DETHERM*, DIOGENES, DRUGU, EMBASE, GMELIN*, HODOC*, HSDB*,
       IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK*, MSDS-OHS, NAPRALERT,
       NIOSHTIC, PDLCOM*, PHAR, PIRA, PROMT, RTECS*, SPECINFO, SYNTHLINE,
       TOXCENTER, ULIDAT, USAN, USPAT2, USPATFULL, VETU
         (*File contains numerically searchable property data)
                      DSL**, EINECS**, TSCA**
     Other Sources:
         (**Enter CHEMLIST File for up-to-date regulatory information)
```

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

7888 REFERENCES IN FILE CA (1957 TO DATE)
503 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
7905 REFERENCES IN FILE CAPLUS (1957 TO DATE)
10 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

139:12012 REFERENCE 1: REFERENCE 139:6868 REFERENCE 139:6791 REFERENCE 139:6446 REFERENCE 139:3078 REFERENCE 138:411210 REFERENCE 7: 138:409366 REFERENCE 8: 138:406796

REFERENCE 9: 138:401298 REFERENCE 10: 138:398399

(CA INDEX

```
L72 ANSWER 12 OF 14 REGISTRY COPYRIGHT 2003 ACS
     110-60-1 REGISTRY
     1,4-Butanediamine (8CI, 9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
     Tetramethylenediamine (7CI)
OTHER NAMES:
CN
     .alpha.,.omega.-Butanediamine
     1,4-Butylenediamine
CN
CN
     1,4-Diamino-n-butane
CN
     1,4-Diaminobutane
CN
     1,4-Tetramethylenediamine
CN
     Putrescin
CN
     Putrescine
FS
     3D CONCORD
MF
     C4 H12 N2
CI
     COM
LC
     STN Files:
                  AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS,
       BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN,
       CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM*,
       DRUGNL, DRUGPAT, DRUGU, DRUGUPDATES, EMBASE, GMELIN*, HODOC*, IFICDB,
       IFIPAT, IFIUDB, IPA, MEDLINE, MRCK*, MSDS-OHS, NAPRALERT, NIOSHTIC,
       PIRA, PROMT, RTECS*, SPECINFO, SYNTHLINE, TOXCENTER, USPAT2, USPATFULL
         (*File contains numerically searchable property data)
     Other Sources:
                      EINECS**, NDSL**, TSCA**
         (**Enter CHEMLIST File for up-to-date regulatory information)
H_2N - (CH_2)_4 - NH_2
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
           10660 REFERENCES IN FILE CA (1957 TO DATE)
             425 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
           10670 REFERENCES IN FILE CAPLUS (1957 TO DATE)
               1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)
            1: 139:7684
REFERENCE
                139:5929
REFERENCE
            2:
               139:3614
REFERENCE
            3:
                139:3528
REFERENCE
                139:2339
REFERENCE
            5:
REFERENCE
            6:
                139:2338
REFERENCE
            7:
                138:411107
REFERENCE
            8:
                138:411018
REFERENCE
            9:
                138:406735
REFERENCE
           10:
                138:406609
```

L72 ANSWER 13 OF 14 REGISTRY COPYRIGHT 2003 ACS

Benzenesulfonamide, N-[(butylamino)carbonyl]-4-methyl- (9CI)

64-77-7 REGISTRY

RN

CN

NAME)

```
OTHER CA INDEX NAMES:
     Urea, 1-butyl-3-(p-tolylsulfonyl)- (8CI)
OTHER NAMES:
     1-Butyl-3-(p-methylphenylsulfonyl)urea
CN
CN
     1-Butyl-3-(p-tolylsulfonyl)urea
CN
     3-(p-Tolyl-4-sulfonyl)-1-butylurea
CN
     Aglicid
CN
     Arkozal
CN
     Artosin
CN
     Artozin
CN
     Butamid
CN
     Butamide
CN
     D 860
CN
     Diaben
CN
     Diabetamid
CN
     Diabetol
CN
     Diabuton
CN
     Diasulfon
CN
     Dolipol
CN
     Glyconon
CN
     HLS 831
     Ipoglicone
CN
CN
     Mobenol
     N-(4-Methylbenzenesulfonyl)-N'-butylurea
CN
     N-(4-Methylphenylsulfonyl)-N'-butylurea
CN
     N-(p-Methylbenzenesulfonyl)-N'-butylurea
CN
     N-(p-Tolylsulfonyl)-N'-butylcarbamide
CN
CN
     N-(Sulfonyl-p-methylbenzene)-N'-n-butylurea
CN
     N-Butyl-N'-(4-methylphenylsulfonyl)urea
CN
     N-Butyl-N'-(p-tolylsulfonyl)urea
     N-Butyl-N'-p-toluenesulfonylurea
CN
     N-n-Butyl-N'-tosylurea
CN
CN
     Orabet
CN
     Oralin
CN
     Orezan
CN
     Orinase
CN
     Orinaz
CN
     Oterben
CN
     Pramidex
CN
     Rastinon
CN
     Tolbusal ·
CN
     Tolbutamid
CN
     Tolbutamide
CN
     Toluina
CN
     Tolumid
CN
     Tolumide
CN
     Toluvan
CN
    U 2043
CN
     Willbutamide
FS
     3D CONCORD
DŔ
     100735-34-0
MF
     C12 H18 N2 O3 S
CI
     COM
LC
     STN Files:
                  ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS,
       BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CEN, CHEMCATS,
       CHEMLIST, CIN, CSCHEM, DDFU, DIOGENES, DRUGU, EMBASE, HODOC*, HSDB*,
       IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK*, NIOSHTIC, PROMT, RTECS*,
       SPECINFO, TOXCENTER, USAN, USPAT2, USPATFULL
         (*File contains numerically searchable property data)
                       DSL**, EINECS**, TSCA**, WHO
         (**Enter CHEMLIST File for up-to-date regulatory information)
```

```
0 0 ||
n-BuNH-C-NH-S ||
0 ||
Me
```

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

3574 REFERENCES IN FILE CA (1957 TO DATE)

25 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

3577 REFERENCES IN FILE CAPLUS (1957 TO DATE)

74 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

REFERENCE 1: 139:294

REFERENCE 2: 138:396425

REFERENCE 3: 138:395332

REFERENCE 4: 138:378974

REFERENCE 5: 138:378518

REFERENCE 6: 138:378502

REFERENCE 7: 138:378464

REFERENCE 8: 138:378406

REFERENCE 9: 138:374137

REFERENCE 10: 138:368761

L72 ANSWER 14 OF 14 REGISTRY COPYRIGHT 2003 ACS

RN **51-85-4** REGISTRY

CN Ethanamine, 2,2'-dithiobis- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Ethylamine, 2,2'-dithiobis- (8CI)

OTHER NAMES:

CN .beta.,.beta.'-Diaminodiethyl disulfide

CN .beta.-Mercaptoethylamine disulfide

CN 1,6-Diamino-3,4-dithiahexane

CN 2,2'-Dithiobis[ethanamine]

CN 2,2'-Dithiobis[ethylamine]

CN 2,2'-Dithiodiethylamine

CN 2-Aminoethane disulfide

CN 2-Aminoethyl disulfide

CN Bis(.beta.-aminoethyl) disulfide

CN Bis(2-aminoethyl) disulfide

CN Cystamine

CN Cysteinamine disulfide

CN Cystineamine

CN Decarboxycystine

CN L 1591

CN Mercamine disulfide

CN Merkamine disulfide

FS 3D CONCORD

MF C4 H12 N2 S2

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, DDFU, DRUGU, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK*, NAPRALERT, NIOSHTIC, PIRA, PROMT, RTECS*, TOXCENTER, USPAT2, USPATFULL

(*File contains numerically searchable property data)
Other Sources: EINECS**

(**Enter CHEMLIST File for up-to-date regulatory information)

H2N-CH2-CH2-S-S-CH2-CH2-NH2

دور سنڌ ج

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1699 REFERENCES IN FILE CA (1957 TO DATE)

62 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

1702 REFERENCES IN FILE CAPLUS (1957 TO DATE) 25 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

REFERENCE 1: 139:13807

REFERENCE 2: 138:395986

REFERENCE 3: 138:384238

REFERENCE 4: 138:338123

REFERENCE 5: 138:268299

REFERENCE 6: 138:216948

REFERENCE 7: 138:203675

REFERENCE 8: 138:188004

REFERENCE 9: 138:175678

REFERENCE 10: 138:140904

=> fil biosis

FILE 'BIOSIS' ENTERED AT 16:14:00 ON 01 JUL 2003 COPYRIGHT (C) 2003 BIOLOGICAL ABSTRACTS INC.(R)

FILE COVERS 1969 TO DATE.

CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 25 June 2003 (20030625/ED)

=> d all tot 182

L82 ANSWER 1 OF 7 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.

AN 1998:120800 BIOSIS

DN PREV199800120800

TI Suppression of aggregate formation and apoptosis by transglutaminase inhibitors in cells expressing truncated DRPLA protein with an expanded polyglutamine stretch.

AU Igarashi, Shuichi; Koide, Reiji (1); Shimohata, Takayoshi; Yamada, Mitsunori; Hayashi, Yasuko; Takano, Hiroki; Date, Hidetoshi; Oyake,

CS

SO

DT

LA

AΒ

ΙT

TT

ΙT

IT

RN

L82 ΑN

DN

TΙ

ΑU

CS

SO

DT

LA

AΒ

```
Mutsuo; Sato, Toshiya; Sato, Aki; Egawa, Shigekimi; Ikeuchi, Takeshi;
     Tanaka, Hajime; Nakano, Ryoichi; Tanaka, Keiko; Hozumi, Isao; Inuzuka,
     Takashi; Takahashi, Hitoshi; Tsuji, Shoji
     (1) Dep. Neurol., Niigatta Univ., 1-757 Asahimachi Niigata 951 Japan
     Nature Genetics, (Feb., 1998) Vol. 18, No. 2, pp. 111-117.
     ISSN: 1061-4036.
     Article
     English
     To elucidate the molecular mechanisms whereby expanded polyglutamine
     stretches elicit a gain of toxic function, we expressed full-length and
     truncated DRPLA (dentatorubral-pallidoluysian
     atrophy) cDNAs with or without expanded CAG repeats in
     COS-7 cells. We found that truncated DRPLA proteins containing an expanded
     polyglutamine stretch form filamentous peri- and intranuclear aggregates
     and undergo apoptosis. The apoptotic cell death was partially suppressed
     by the transglutaminase inhibitors cystamine and
     monodansyl cadaverine (but not putrescine), suggesting
     involvement of a transqlutaminase reaction and providing a
     potential basis for the development of therapeutic measures for
     CAG-repeat expansion diseases.
     Genetics and Cytogenetics - Animal *03506
     Cytology and Cytochemistry - Animal *02506
     Biochemical Studies - Proteins, Peptides and Amino Acids
     Enzymes - Chemical and Physical *10806
     Pathology, General and Miscellaneous - General *12502
     Pathology, General and Miscellaneous - Necrosis *12510
BC ·
     Cercopithecidae
                       86205
     Major Concepts
        Molecular Genetics (Biochemistry and Molecular Biophysics)
     Diseases
          CAG-repeat expansion diseases: genetic disease
     Chemicals & Biochemicals
        cDNA [complementary DNA]; transglutaminase; DRPLA protein [
        dentatorubral-pallidoluysian atrophy
        protein]: expanded polyglutamine stretch, toxic function
     Miscellaneous Descriptors
        aggregate formation; apoptotic cell death
ORGN Super Taxa
        Cercopithecidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        COS-7 (Cercopithecidae)
ORGN Organism Superterms
        Animals; Chordates; Mammals; Nonhuman Mammals; Nonhuman Primates;
        Nonhuman Vertebrates; Primates; Vertebrates
     80146-85-6 (TRANSGLUTAMINASE)
       26700-71-0Q (POLYGLUTAMINE)
       69864-43-3Q (POLYGLUTAMINE)
     ANSWER 2 OF 7 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.
     1997:300704 BIOSIS
     PREV199799599907
     Pharmacologic inhibition of transglutaminase-induced
     cross-linking of Alzheimer's amyloid beta-peptide.
     Zhang, Wei; Johnson, Brett R.; Bjornsson, Thorir D. (1)
     (1) Div. Clin. Pharmacol., Dep. Med., Jefferson Med. Coll., 1100 Walnut
     St., MOB-601, Philadelphia, PA 19107 USA
     Life Sciences, (1997) Vol. 60, No. 25, pp. 2323-2332.
     ISSN: 0024-3205.
     Article
     English
     The brain of Alzheimer's disease (AD) patients contains deposits of
     amyloid beta-peptide (A-beta). Recent studies have shown that A-beta is a
```

substrate for tissue transglutaminase (TGase), which induces the

```
formation of cross-linked dimers and polymers, and that tacrine,
     indomethacin and deferoxamine, which have widely different chemical
     structures, attenuate the progression of symptoms of AD. This report
     evaluated the potential of a total of ten different pharmacological agents
     to inhibit TGase-induced cross-linking of A-beta, including known TGase
     inhibitors (dansylcadaverine, spermine), non-steroidal
     anti-inflammatory drugs (indomethacin, meclofenamic acid, diflunisal,
     salicylic acid), monoamine oxidase inhibitors (tranylcypromine,
     phenelzine), an acetylcholinesterase inhibitor (tacrine), and an iron
     chelating agent (deferoxamine). All but one (salicylic acid) of these ten
     agents had an inhibitory effect on TGase-induced A-beta cross-linking.
     These results suggest that inhibition of TGase-induced cross-linking of
     A-beta is a potential pharmacologic target for the treatment of AD. A
     method is also presented for the determination of percent inhibition of
     TGase-induced A-beta cross-linking based on the separated monomer, dimer
     and polymer bands on SDS-PAGE gels.
     Biochemical Studies - General *10060
     Biochemical Studies - Proteins, Peptides and Amino Acids *10064
     Biophysics - Molecular Properties and Macromolecules *10506
     Enzymes - Chemical and Physical
       Pharmacology - Neuropharmacology *22024
     Major Concepts
        Biochemistry and Molecular Biophysics; Enzymology (Biochemistry and
        Molecular Biophysics); Pharmacology
     Chemicals & Biochemicals
          TRANSGLUTAMINASE; AMYLOID; DANSYLCADAVERINE;
        SPERMINE; INDOMETHACIN; MECLOFENAMIC ACID; DIFLUNISAL; SALICYLIC ACID;
        TRANYLCYPROMINE; PHENELZINE; TACRINE; DEFEROXAMINE; EC 2.3.2.13
     Miscellaneous Descriptors
        ALZHEIMER'S DISEASE; AMYLOID BETA-PEPTIDE; ANTI-ALZHEIMER'S AGENT;
        ANTIINFLAMMATORY-DRUG; BIOCHEMISTRY AND BIOPHYSICS;
       DANSYLCADAVERINE; DEFEROXAMINE; DIFLUNISAL; EC 2.3.2.13;
        INDOMETHACIN; MECLOFENAMIC ACID; NERVOUS SYSTEM DISEASE; PHARMACOLOGY;
       PHENELZINE; SALICYLIC ACID; SPERMINE; TACRINE; TRANSGLUTAMINASE
        ; TRANSGLUTAMINASE INHIBITOR; TRANSGLUTAMINASE
        -INDUCED CROSS-LINKING INHIBITION; TRANYLCYPROMINE
     80146-85-6 (TRANSGLUTAMINASE)
     11061-24-8 (AMYLOID)
       10121-91-2 (DANSYLCADAVERINE)
     71-44-3 (SPERMINE)
     53-86-1 (INDOMETHACIN)
     644-62-2 (MECLOFENAMIC ACID)
     22494-42-4 (DIFLUNISAL)
     69-72-7 (SALICYLIC ACID)
     155-09-9 (TRANYLCYPROMINE)
     51-71-8 (PHENELZINE)
     321-64-2 (TACRINE)
     70-51-9 (DEFEROXAMINE)
       80146-85-6 (EC 2.3.2.13)
     ANSWER 3 OF 7 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.
L82
     1997:113658 BIOSIS
     PREV199799412861
     Microtubules and microfilaments participate in the inhibition of
     synaptosomal noradrenaline release by tetanus toxin.
     Ashton, Anthony C.; Dolly, J. Oliver (1)
     (1) Dep. Biochemistry, Imperial Coll., London SW7 2AY UK
     Journal of Neurochemistry, (1997) Vol. 68, No. 2, pp. 649-658.
     ISSN: 0022-3042.
       ticle
```

√s toxin (TeTX) has been demonstrated to inhibit transmitter release o mechanisms: Zn-2+dependent proteolytic cleavage of synaptobrevin

CC

ΙT

ΙT

IΤ

RN

AN

DN

ΤI

ΑU

CS

and activation of a neuronal transglutaminase. Herein, attenuation of TeTX-induced blockade of noradrenaline release from synaptosomes was achieved by prior disassembly of microfilaments with cytochalasin D or breakdown of microtubules by colchicine or nocodazole. These drugs and monodansylcadaverine, a transglutaminase inhibitor, displayed some additivity in antagonizing the inhibitory effect of the toxin on synaptosomal transmitter release; as none of them reduced synaptobrevin cleavage, all appear to work independently of the toxin's proteolytic action. Prior stabilization of microtubules with taxol prevented the antagonism seen with colchicine, highlighting that this cytoskeletal component is the locus of the effect of colchicine. Replacement of Ca-2+ with Ba-2+ caused disappearance of the fraction of evoked secretion whose inhibition by TeTX is reliant on polymerized actin but did not alter the blockade by toxin that is dependent on microtubules. Two temporally distinguished phases of release were reduced by TeTX, and colchicine lessened its effects on both. Blockade of the fast phase (ltoreq 10 s) of secretion by TeTX was unaffected by cytochalasin D, but it clearly antagonized the toxin-induced inhibition of the slow (10-s to gtoreg 5-min) component; it is notable that such antagonism was accentuated during a second bout of evoked release. These findings are consistent with sustained release requiring dissociation of synaptic vesicles from the microfilaments, a step that seems to be perturbed by TeTX.

CC Cytology and Cytochemistry - Animal *02506 Enzymes - Physiological Studies *10808 Metabolism - Proteins, Peptides and Amino Acids *13012 Endocrine System - Neuroendocrinology *17020

Nervous System - Pathology *20506

Toxicology - General; Methods and Experimental *22501 Physiology and Biochemistry of Bacteria *31000

BC Muridae *86375

IT Major Concepts

Cell Biology; Endocrine System (Chemical Coordination and Homeostasis); Enzymology (Biochemistry and Molecular Biophysics); Metabolism; Nervous System (Neural Coordination); Physiology; Toxicology

IT Chemicals & Biochemicals

NORADRENALINE; NOREPINEPHRINE; TRANSGLUTAMINASE

IT Miscellaneous Descriptors

MICROFILAMENT; MICROTUBULE; NERVOUS SYSTEM; NOREPINEPHRINE; SYNAPTIC VESICLE; SYNAPTOBREVIN; SYNAPTOSOMAL RELEASE; TETANUS TOXIN;

TRANSGLUTAMINASE

ORGN Super Taxa

Muridae: Rodentia, Mammalia, Vertebrata, Chordata, Animalia

ORGN Organism Name

rat (Muridae)

ORGN Organism Superterms

animals; chordates; mammals; nonhuman mammals; nonhuman vertebrates; rodents; vertebrates

RN 51-41-2 (NORADRENALINE)

51-41-2 (NOREPINEPHRINE)

80146-85-6 (TRANSGLUTAMINASE)

- L82 ANSWER 4 OF 7 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC.
- AN 1996:416490 BIOSIS
- DN PREV199699138846
- TI Role of **transglutaminase** in (3H)5-HT release from synaptosomes and in the inhibitory effect of tetanus toxin.
- AU Gobbi, M. (1); Frittoli, E.; Mennini, T.
- CS (1) Istituto di Ricerche Farmacologiche "Mario Negri", Via Eritrea 62, 20157 Milano Italy
- SO Neurochemistry International, (1996) Vol. 29, No. 2, pp. 129-134. ISSN: 0197-0186.
- DT Article

LA English AΒ It has been suggested that the Ca-2+-dependent enzyme transglutaminase (TGase) may suppress vesicular neurotransmitter release and mediate the inhibitory effect of tetanus toxin (TetTx) on exocytosis. The aim of the present study was to test this in a model of (3H)5-HT release from superfused rat cortical synaptosomes. Monodansylcadaverine, a synthetic amine chat acts as an alternative substrate for TGase, showed dose-dependent releasing activity which, however, was Ca-2+-independent, being maintained in a Ca-2+-free buffer (containing EGTA) or using synaptosomes preloaded with the intracellular Ca-2+ chelator BAPTA. Preincubation of synaptosomes with RS-48373, an irreversible TGase inactivator, resulted in marked (64%) and persistent inhibition of endogenous TGase but did not alter basal and K+-induced (3H)5-HT release. Preincubation of synaptosomes with 10 nM TetTx resulted in 52% inhibition of K+-induced (3H)5-HT release, and this effect was not antagonized in RS-48373-treated synaptosomes. The inhibitory effect of TetTx was significantly antagonized by 20 mM captopril, a metalloendoprotease inhibitor, confirming in rat brain synaptosomes that TetTx inhibits exocytosis by acting as a metalloendoprotease. These results suggest that TGase is not involved in controlling (3H)5-HT release from resting and depolarized synaptosomes, or in the inhibitory effect of TetTx. Biochemical Studies - Proteins, Peptides and Amino Acids Biochemical Studies - Minerals 10069 Enzymes - Physiological Studies *10808 *17020 Endocrine System - Neuroendocrinology Muscle - Pathology *17506 Nervous System - Physiology and Biochemistry *20504 Nervous System - Pathology *20506 Toxicology - General; Methods and Experimental *86375 BCMuridae Major Concepts ΙT Endocrine System (Chemical Coordination and Homeostasis); Enzymology (Biochemistry and Molecular Biophysics); Muscular System (Movement and Support); Nervous System (Neural Coordination); Toxicology IT Chemicals & Biochemicals TRANSGLUTAMINASE; CALCIUM (II); POTASSIUM (I); SEROTONIN Miscellaneous Descriptors IT CALCIUM (II); CORTEX; POTASSIUM (I); SEROTONIN ORGN Super Taxa Muridae: Rodentia, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name rat (Muridae) ORGN Organism Superterms animals; chordates; mammals; nonhuman mammals; nonhuman vertebrates; rodents; vertebrates 80146-85-6 (TRANSGLUTAMINASE) RN 14127-61-8 (CALCIUM (II)) 24203-36-9 (POTASSIUM (I)) 50-67-9 (SEROTONIN) ANSWER 5 OF 7 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. L82 1987:29797 BIOSIS ANBA83:19731 DN ΤI CYTOTOXIC EFFECTS OF MONODANSYLCADAVERINE AND METHYLAMINE IN PRIMARY CULTURES OF RAT CEREBELLAR NEURONS. ΑU GILAD G M; GILAD V H VA MED. CENT., SPINAL CORD INJURY RES., 1400 VETERANS OF FOREIGN WARS CS PARKWAY, BOSTON, MASS. 02132, USA. INT J DEV NEUROSCI, (1986) 4 (5), 401-406. SO CODEN: IJDND6. ISSN: 0736-5748.

FS

LA

BA; OLD .

English

AB The effects of dansylcadaverine and methylamine, competitive inhibitors of transglutaminase, were examined in primary cultures of dissociated rat cerebellar neurons. Addition of the drugs at plating time resulted 24 hr later in irreversible cytotoxic effects evidenced by failure of aggregation and neurite formation. Cytotoxicity was dose-dependent with methylamine being more potent (IC50 = 20 .mu.M) than dansylcadaverine (IC50 = 30 .mu.M). The cytotoxic effects were less potent when drugs were added 24 hr after plating, the time when neurons had already begun to extend neurites. Drugs were effective in the various sera and heat-inactivated sera tested. We concluded that low doses of methylamine and dansylcadaverine have potent toxic effects on primary neuronal cultures. Cytology and Cytochemistry - Animal *02506 Enzymes - Physiological Studies *10808

Pathology, General and Miscellaneous - Comparative Pathology, General and Miscellaneous - Necrosis

Nervous System - Pathology *20506

Toxicology - Pharmacological Toxicology Tissue Culture, Apparatus, Methods and Media 32500 In Vitro Studies, Cellular and Subcellular 32600 Plant Physiology, Biochemistry and Biophysics - Chemical Constituents 51522

Pharmacognosy and Pharmaceutical Botany 54000

BC Bovidae 85715 Hominidae 86215

Muridae 86375 ΙT

Miscellaneous Descriptors FETAL CALF HUMAN TRANSGLUTAMINASE INHIBITORS

RN 74-89-5 (METHYLAMINE)

10121-91-2 (MONODANSYLCADAVERINE) 80146-85-6 (TRANSGLUTAMINASE)

- ANSWER 6 OF 7 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. L82
- 1983:209038 BIOSIS AN
- DN BA75:59038
- BRAIN TRANS GLUTAMINASE EC-2.3.2.13 IN-VITRO CROSS TΙ LINKING OF HUMAN NEURO FILAMENT PROTEINS INTO INSOLUBLE POLYMERS.
- AU SELKOE D J; ABRAHAM C; IHARA Y
- CS RALPH LOWELL LABORATOIRES, MAILMAN RESEARCH CENT., MCLEAN HOSP., BELMONT, MASS. 02178.
- SO PROC NATL ACAD SCI U S A, (1982) 79 (19), 6070-6074. CODEN: PNASA6. ISSN: 0027-8424.
- FS BA; OLD
- English T.A
- The accumulation in aged human neurons of insoluble, high-MW filamentous AB polymers apparently linked by nondisulfide covalent bonds led to the examination of human brain for the presence of transglutaminase (EC 2.3.2.13) and endogenous protein substrates for this crosslinking enzyme. The presence of a transamidating enzyme that can covalently crosslink brain proteins into insoluble polymers in vitro by forming .gamma.-glutamyl-.epsilon.-lysine intermolecular bridges was demonstrated. Brain transglutaminase is Ca2+ dependent, has an electrophoretic mobility similar to that of erythrocyte transglutaminase and is active in human postmortem brain from aged normal individuals and patients with Alzheimer disease (senile dementia). Brain neurofilament fractions incubated in the presence of transglutaminase, Ca2+ and the fluorescent amine dansylcadaverine form a fluorescent, nondisulfide-bonded insoluble polymer; this process is associated with a decrease in the amount of soluble neurofilament polypeptides in the preparation. EM of the polymeric material reveals an extensive network of connecting filaments, which can be immunostained with various neurofilament antisera. Cystamine, an inhibitor of transglutaminase, prevents the neurofilament crosslinking. Glial

filaments and myelin basic protein can also serve as substrates of brain transglutaminase in vitro. Although Alzheimer disease-type paired helical filaments were not formed under the specific in vitro coinditions employed, the data suggest 1 possible mechanism for the covalent crosslinking of filaments into insoluble polymers during human neuronal aging. Microscopy Techniques - Electron Microscopy 01058 Biochemical Studies - Proteins, Peptides and Amino Acids 10064 Biochemical Studies - Lipids 10066 Biochemical Studies - Minerals 10069 Biophysics - Molecular Properties and Macromolecules 10506 Enzymes - Physiological Studies *10808 Anatomy and Histology, General and Comparative - Microscopic and Ultramicroscopic Anatomy *11108 Metabolism - Proteins, Peptides and Amino Acids Nervous System - Physiology and Biochemistry *20504 Nervous System - Pathology *20506 Psychiatry - Psychopathology; Psychodynamics and Therapy *21002 Gerontology *24500 Developmental Biology - Embryology - Morphogenesis, General *25508 Hominidae 86215 Miscellaneous Descriptors ELECTRON MICROSCOPY CYSTAMINE DANSYL CADAVERINE ALZHEIMER DISEASE AGING MYELIN BASIC PROTEIN NEURONAL AGING CALCIUM 51-85-4 (CYSTAMINE) 7440-70-2 (CALCIUM) 10121-91-2 (DANSYL CADAVERINE) 80146-85-6 (EC-2.3.2.13) 80146-85-6 (TRANS GLUTAMINASE) L82 ANSWER 7 OF 7 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. 1983:79120 BIOSIS BR25:4120 REGULATION OF ORNITHINE DECARBOXYLASE EC-4.1.1.17 ACTIVITY IN CULTURED GLIOMA CELLS BY TRANS GLUTAMINASE EC-2.3.2.13. KORNER G; BACHRACH U DEP. MOL. BIOL., HEBREW UNIV.-HADASSAH MED. SCH., JERUSALEM. THE 1982 ANNUAL MEETING OF THE ISRAEL BIOCHEMICAL SOCIETY IN CONJUNCTION WITH THE SECTION FOR BIOTECHNOLOGY AND THE ISRAEL BIOPHYSICAL SOCIETY, REHOVOT, ISRAEL, APRIL 11-12, 1982. ISR J MED SCI. (1982) 18 (6), 11. CODEN: IJMDAI. ISSN: 0021-2180. Conference BR; OLD English General Biology - Symposia, Transactions and Proceedings of Conferences, Congresses, Review Annuals 00520 Cytology and Cytochemistry - Animal 02506 Biochemical Studies - Proteins, Peptides and Amino Acids 10064 Enzymes - Chemical and Physical Enzymes - Physiological Studies *10808 Metabolism - Proteins, Peptides and Amino Acids *13012 Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies 15002 Nervous System - Physiology and Biochemistry *20504 Nervous System - Pathology 20506 Neoplasms and Neoplastic Agents - Neoplastic Cell Lines 24005 Developmental Biology - Embryology - Morphogenesis, General *25508 Tissue Culture, Apparatus, Methods and Media 32500 Bovidae 85715 Muridae 86375 Miscellaneous Descriptors

ABSTRACT RAT C-6 BU-1 CELLS PUTRESCINE ISOPROTERENOL FETAL CALF SERUM

DANSYL CADAVERINE GROWTH DIFFERENTIATION

BC

ΙT

RN

ΑN

DN

TI

ΑU

CS

SO

DT

FS

LA

CC

BC

ΙT

RN 110-60-1 (PUTRESCINE) 7683-59-2 (ISOPROTERENOL) 9024-60-6 (ORNITHINE DECARBOXYLASE EC-4.1.1.17) 10121-91-2 (DANSYL CADAVERINE) **80146-85-6** (EC-2.3.2.13) 80146-85-6 (TRANS GLUTAMINASE) => fil wpix FILE 'WPIX' ENTERED AT 16:21:20 ON 01 JUL 2003 COPYRIGHT (C) 2003 THOMSON DERWENT 30 JUN 2003 FILE LAST UPDATED: <20030630/UP> 200341 MOST RECENT DERWENT UPDATE: <200341/DW> DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE >>> NEW WEEKLY SDI FREQUENCY AVAILABLE --> see NEWS < >>> ${\tt SLART}$ (Simultaneous Left and Right Truncation) is now available in the /ABEX field. An additional search field /BIX is also provided which comprises both /BI and /ABEX <<< >>> PATENT IMAGES AVAILABLE FOR PRINT AND DISPLAY <<< >>> FOR DETAILS OF THE PATENTS COVERED IN CURRENT UPDATES, SEE http://www.derwent.com/dwpi/updates/dwpicov/index.html <<< >>> FOR A COPY OF THE DERWENT WORLD PATENTS INDEX STN USER GUIDE, PLEASE VISIT: http://www.stn-international.de/training center/patents/stn guide.pdf <<< >>> FOR INFORMATION ON ALL DERWENT WORLD PATENTS INDEX USER GUIDES, PLEASE VISIT: http://www.derwent.com/userguides/dwpi guide.html <<< => d 191 all abeq tech abex tot L91 ANSWER 1 OF 3 WPIX (C) 2003 THOMSON DERWENT AN - 2000-147134 [13] WPIX DNC C2000-046006 Treatment of neurodegenerative diseases and other diseases mediated by an enzyme activity, e.g. Huntington's disease. DC · B04 B05 KARPUJ, M V; STEINMAN, L IN (YEDA) YEDA RES & DEV CO LTD PA CYC 86 A1 19991223 (200013) * EN WO 9965516 61p A61K038-48 PΙ RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ UG ZW W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG US UZ VN YU ZA ZW A 20000105 (200024) A61K038-48 AU 9948239 WO 9965516 A1 WO 1999-US13615 19990617; AU 9948239 A AU 1999-48239 ADT 19990617 FDT AU 9948239 A Based on WO 9965516 19980617 PRAI US 1998-89603P ICM A61K038-48 IC ICS A61K031-13 AΒ 9965516 A UPAB: 20000313 NOVELTY - Treatment of transglutaminase (T) mediated diseases

involves administration of its inhibitor (TI).

ACTIVITY - Nootropic; antirheumatic; neuroprotective; antidiabetic; antiinflammatory. The antiinflammatory activity of a (TI), monodansyl cadaverine was tested using paraparetic mice with experimental autoimmune encephalomyelitis induced by injecting 4 mg of mouse spinal cord homogenate. 0.05 mM monodansyl cadaverine was injected intraperitoneally into one of two groups of mice after 13 days of disease induction. A significant influence (p=0.03 compared to control) of this inhibitor occured after the second day of injection and mice treated with the monodansyl cadaverine showed reversal in the paralytic disease.

MECHANISM OF ACTION - Transglutaminase inhibitor.

USE - The composition comprising (TI) is used in the treatment of (TI) mediated diseases like neurodegenerative diseases caused by aggregation of polyQ proteins, Huntington's disease, spinobulbar atrophy, spinocerebellar ataxia, dentatorubralpallidoluysian atrophy, cell mediated autoimmune disease like rheumatoid arthritis, multiple sclerosis or insulin dependent diabetes mellitus and other inflammatory diseases of the central nervous system (claimed).

ADVANTAGE - None given.

Dwg.0/7

FS CPI

FA AB; DCN

MC CPI: B04-E02; B04-E06; B04-F0400E; B04-F1100E; B14-C03; B14-C06;

B14-D06; B14-E08; B14-J01; B14-J01A;

B14-J01A4; B14-S03; B14-S04

TECH

UPTX: 20000313

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Method: (TI) can also be presented as an antisense DNA of (T) gene or as DNA encoding it and is introduced into the cell of patient. The method of DNA introduction includes receptor mediated gene delivery, transkaryotic implantation, viral shuttle vectors, direct injection of non-infectious, non-oncogenic plasma DNA encapsulated in liposomes, immunoliposomes and a liposome/red blood cell membrane hybrid.

ABEX

UPTX: 20000313

SPECIFIC COMPOUNDS - The specific (TI) compounds are monodansyl cadaverine, cystamine, putrescine, gamma-amino benzoic acid, N-benzyloxy carbonyl, 5-deazo-4-oxonorvaline p-nitrophenylester, glycine methyl ester, CuSO4, and tolbutamide (claimed).

ADMINISTRATION - Administration can be by any preferred route e.g. intraperitoneal, subcutaneous, oral routes and are given in dosages of 0.0001-100 mg/kg body weight daily.

L91 ANSWER 2 OF 3 WPIX (C) 2003 THOMSON DERWENT

AN 1998-130408 [12] WPIX

DNC C1998-043052

TI Use of modulators of trans glutaminase activity - in promoting healing of wounds, chronic wounds and fibrotic disorders with reduced scarring.

DC B04 B05 D16

IN FERGUSON, M W J

PA (UYMA-N) UNIV VICTORIA MANCHESTER

CYC 72

PI WO 9804245 A1 19980205 (199812) * EN 30p A61K031-00

RW: AT BE CH DE DK EA ES FI FR GB GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG

W: AL AM AT AU AZ BB BG BR BY CA CH CN CZ DE DK EE ES FI GB GE HU IL IS JP KE KG KP KR KZ LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK TJ TM TR TT UA UG US UZ VN

ZA 9606486 A 19980429 (199822)# 28p C12N000-00

AU 9666205 A 19980220 (199828) A61K031-00

ADT WO 9804245 A1 WO 1996-GB1785 19960725; ZA 9606486 A ZA 1996-6486 19960731; AU 9666205 A AU 1996-66205 19960725, WO 1996-GB1785 19960725

```
FDT AU 9666205 A Based on WO 9804245
PRAI WO 1996-GB1785
                      19960725; ZA 1996-6486
                                                 19960731
     ICM A61K031-00; C12N000-00
          A61K031-13; A61K031-145; A61K031-18; A61K031-40; A61K038-12;
          C07C000-00
          9804245 A UPAB: 19980323
ΑB
     Use of an inhibitor of transglutaminase in promoting healing of
     wounds or fibrotic disorders with reduced scarring.
          Also claimed is the use of a stimulator of transglutaminase
     activity in promoting healing of chronic wounds.
          The transglutaminase is type II transglutaminase.
     The inhibitor is an active site inhibitor or is a substrate competitive
     inhibitor. It is selected from cystamine, monodansyl
     cadaverine, 2-(3-(diallylamino) propionyl) benzothiophene,
     putrescine, bacitracin, a neutralising antibody (or an antigen-binding
     fragment of this specific to transglutaminase) or a derivative
     of 2((2-oxopropyl)thio) imidazolium. The neutralising antibody is an IgG
     antibody specific to transglutaminase and is a monoclonal
     antibody, polyclonal antibody or genetically engineered antibody.
          USE - The transglutaminase inhibitor is used for promoting
     healing of wounds (e.g. skin wounds, tendon damage, crush injuries, eye
     wounds, central nervous system injuries or scar tissue formation resulting
     from strokes) or fibrotic disorders (e.g. pulmonary fibrosis,
     glomerulonephritis, liver cirrhosis or proliferative vitreo retinopathy).
     The transglutaminase stimulator may be used in treatment of
     chronic wounds (e.g. venous ulcers, diabetic ulcers or bed sores).
          ADVANTAGE - The inhibitor/stimulator reduce scarring. Scar tissue
     formation provides mechanical strength to healed wounds but can be
     unsightly and may impair the function of the tissue. This is especially
     the case in the central nervous system.
     Dwg.0/0
     CPI
FS
FA
     AB; DCN
MC
     CPI: B04-C01; B04-G01; B04-L05; B06-H; B07-H; B14-N17B; D05-C03
                           (C) 2003 THOMSON DERWENT
     ANSWER 3 OF 3 WPIX
1.91
     1991-207851 [28]
                        WPIX
AN
DNC
    C1991-090106
ΤI
     Growth arrest of eukaryotic cells (e.g. B and T lymphocytes) - with the
     trans-glutaminase inhibitors mono-
     dansyl cadaverine or 1-(5-aminopentyl)-3-(phenyl-
     thiourea) useful as adjunct to chemotherapy.
DC
     MEHTA, K; SAHASRABUDDHE, C G; SAHASRABUD, C G
IN
     (TEXA) UNIV TEXAS SYSTEM
PΑ
CYC
     30
                   A 19910627 (199128)*
PΤ
     WO 9108739
        RW: AT BE CH DE DK ES FR GB GR IT LU NL OA SE
         W: AT AU BB BG CA CH DE DK ES FI GB GR HU JP KR LU MC MG MW NL NO RO
            SD SE SU
     AU 9171618
                     19910718 (199142)
     EP 505487
                   A1 19920930 (199240)
                                         EN
                                              44p
                                                     A61K031-17
         R: AT BE CH DE DK ES FR GB GR IT LI LU NL SE
     JP 05502673
                   W
                     19930513 (199324)
                                              11p
                                                     A61K045-00
     EP 505487 A1 WO 1990-US7349 19901213, EP 1991-902418 19901213; JP 05502673
ADT
     W WO 1990-US7349 19901213, JP 1991-502790 19901213
     EP 505487 Al Based on WO 9108739; JP 05502673 W Based on WO 9108739
PRAI US 1989-451324
                      19891213
REP
     6.Jnl.Ref
     ICM A61K031-17; A61K045-00
IC
     ICS
         A61K031-18
          9108739 A UPAB: 19930928
AB
     The transglutaminase inhibitor is e.g. monodansyl
```

cadaverine (MDC), which may be used to arrest the growth of B
lymphocytes, T lymphocytes or monocytes (in a cell cycle G1-phase).
Another cpd. 1-(5-aminopentyl)-3-phenylthiourea (PPTU), is claimed for arresting B lymphocyte growth.

The inhibitors arrest the growth of normal and neoplastic B and T lymphocytes, preventing their entry into the S-phase of the cell cycle. The in vitro activity of MDC and PPTU indicate that these cpds. would be highly desirable adjunct to chemotherapy.

0/8

FS CPI

FA AB; DCN

MC CPI: B10-A08; B10-A13A; B12-G01B3; B12-G05; B12-G07

ABEQ JP 05502673 W UPAB: 19931116

The transglutaminase inhibitor is e.g. monodansyl cadaverine (MDC), which may be used to arrest the growth of B lymphocytes, T lymphocytes or monocytes (in a cell cycle G1-phase). Another cpd. 1-(5-aminopentyl)-3-phenylthiourea (PPTU), is claimed for arresting B lymphocyte growth.

The inhibitors arrest the growth of normal and neoplastic B and T lymphocytes, preventing their entry into the S-phase of the cell cycle. The in vitro activity of MDC and PPTU indicate that these cpds. would be highly desirable adjunct to chemotherapy.

=> d his

(FILE 'HOME' ENTERED AT 15:24:52 ON 01 JUL 2003) SET COST OFF

FILE 'HCAPLUS' ENTERED AT 15:25:07 ON 01 JUL 2003

E W099-US13615/AP, PRN

L1 1 S E3.E4

E W09965516/PN

L2 1 S E3

L3 1 S L1, L2

FILE 'REGISTRY' ENTERED AT 15:26:23 ON 01 JUL 2003

L4 1 S 80146-85-6

L5 1 S 10121-91-2

L6 0 S 10121-91-2/CRN

FILE 'HCAPLUS' ENTERED AT 15:27:17 ON 01 JUL 2003

L7 265 S L5

 $\Gamma8$

L13

568 S DANSYLCADAVERIN# OR MONODANSYLCADAVERIN# OR (MONODANSYL OR DA

L9 6 S N 5 AMINOPENTYL 5 DIMETHYLAMINO 1 NAPHTHALENESULFONAMIDE

L10 617 S L7-L9

L11 538 S L10 AND (PD<=19980617 OR PRD<=19980617 OR AD<=19980617)

E STEINMAN L/AU

L12 208 S E3, E4

1 S L10 AND L12

L14 1 S YEDA?/PA,CS AND L10

L15 1 S L3, L13, L14

L16 2799 S L4

L17 3813 S TRANSGLUTAMINASE OR TRANS GLUTAMINASE

L18 228. S L11 AND L16, L17

E HUNTINGTON/CT

E E5+ALL

E NERVOUS SYTEM/CT

E NERVOUS SYSTEM/CT

L19 3155 S NERVOUS SYSTEM/CT (L) (HUNTINGTON? OR CHOREA?)

E SPINOBULBAR/CT

E SPINOBULBAR

L20 85 S E2-E6

```
L21
             55 S L20 (L) ATROPH?
                E SPINOCEREBELLAR/CT
                E E4+ALL
L22
            471 S E2
                E DENTATORUBRAL
L23
            256 S E3-E8
                E BRAIN DISEASE/CT
                E E4+ALL
                E E2+ALL
L24
            179 S E3-E5 (L) DENTATORUB?
L25
            179 S E3-E5 (L) PALLIDOL?
L26
              3 S L11 AND L19-L25
              3 S L11 AND (HUNTINGTON? OR CHOREA? OR SPINOBULBAR? OR SPINOCEREB
L27
              3 S L26, L27 AND L18
L28
     FILE 'REGISTRY' ENTERED AT 15:44:40 ON 01 JUL 2003
              5 S 51-85-4 OR 110-60-1 OR 64-77-7 OR 7758-98-7 OR 616-34-2
L29
              2 S 150-13-0 OR 74389-76-7
L30
            393 S 7664-93-9/CRN AND CU/ELS
L31
             47 S L31 AND 2/NC
L32
             44 S L32 NOT (CCS OR MXS OR IDS OR MNS OR PMS)/CI
L33
             11 S L33 NOT ATO
L34
L35
              5 S L34 AND NR>=1
L36
              6 S L34 NOT L35
     FILE 'HCAPLUS' ENTERED AT 15:50:38 ON 01 JUL 2003
          43274 S L29 OR L30 OR L36
L37
           180 S L37 AND L16, L17
L38
L39
             15 S L37 AND L19-L25
             48 S L37 AND (HUNTINGTON? OR CHOREA? OR SPINOBULBAR? OR SPINOCEREB
L40
              8 S L39, L40 AND L38
L41
          37213 S L37 AND (PD<=19980617 OR PRD<=19980617 OR AD<=19980617)
L42
            154 S L42 AND L38
L43
             31 S L42 AND L39, L40
L44
L45
              4 S L43 AND L44
              5 S L28, L45
L46
              3 S L12 AND L37
L47
              7 S L46, L47
L48
L49
             1 S L11, L42 AND (POLYQ OR POLY Q)
     FILE 'REGISTRY' ENTERED AT 15:55:59 ON 01 JUL 2003
L50
              2 S 24991-23-9 OR 25513-46-6
L51
              4 S (5959-95-5 OR 6899-04-3)/CRN AND PMS/CI
              2 S L51 AND 1/NC
L52
              1 S 101985-79-9
L53
                E CAG
L54
            194 S E3
L55
              2 S L54 AND GUAN?
     FILE 'HCAPLUS' ENTERED AT 15:59:40 ON 01 JUL 2003
L56
           2463 S L50, L52, L53
             19 S L56 AND L11, L42
L57
              4 S L57 AND L16, L17
L58
              3 S L58 NOT CLOTTING/TI
L59
L60
              7 S L48, L49, L59
              7 S L60 AND L1-L3, L7-L28, L37-L49, L56-L60
L61
     FILE 'REGISTRY' ENTERED AT 16:02:56 ON 01 JUL 2003
              2 S 26700-71-0 OR 69864-43-3
L62
            118 S (6893-26-1 OR 617-65-2)/CRN AND PMS/CI
L63
L64
              2 S L63 AND 1/NC
```

FILE 'HCAPLUS' ENTERED AT 16:04:25 ON 01 JUL 2003

```
L65
            896 S L62, L64
            386 S L65 AND (PD<=19980617 OR PRD<=19980617 OR AD<=19980617)
L66
              9 S L66 AND L16, L17
L67
L68
             64 S L66 AND L19-L25
              6 S L67 AND L68
L69
L70
             11 S L61, L69
             11 S L70 AND L1-L3, L7-L28, L37-L49, L56-L61, L65-L70
L71
     FILE 'HCAPLUS' ENTERED AT 16:07:58 ON 01 JUL 2003
                SEL HIT RN
     FILE 'REGISTRY' ENTERED AT 16:08:12 ON 01 JUL 2003
             14 S E1-E14
L72
     FILE 'BIOSIS' ENTERED AT 16:08:28 ON 01 JUL 2003
L73
            564 S L10 ·
            470 S L73 AND PY<=1998
L74
L75
            209 S L74 AND L16, L17
              1 S L74 AND (HUNTINGTON? OR CHOREA? OR POLYQ OR POLY Q OR CAG OR
L76
              1 S L74 AND L62, L64, L50, L52, L53
L77
L78
             1 S L76, L77
             1 S L75 AND L78
L79
             16 S L74 AND (20506 OR 22024)/CC
L80
              6 S L75, L79 AND L80
L81
              7 S L79, L81
L82
             10 S L80 NOT L82
L83
     FILE 'BIOSIS' ENTERED AT 16:14:00 ON 01 JUL 2003
     FILE 'WPIX' ENTERED AT 16:14:19 ON 01 JUL 2003
             11 S L8/BIX OR L9/BIX
L84
             E DANSYLCADAVERINE/DCN
                E DANSYL CADAVERINE/DCN
                E CADAVERINE/DCN
                E E3+ALL
                E R21595+ALL/DCN
              5 S L84 AND (TRANSGLUTAMINASE OR TRANS GLUTAMINASE)/BIX
              1 S L84 AND (B14-D06 OR C14-D06 OR B12-G01B1 OR C12-G01B1)/MC
L86
L87
              1 S L84 AND (B14-J? OR C14-J? OR B12-C? OR C12-C?)/MC
              1 S L84 AND (B14-S03? OR C14-S03?)/MC
L88
L89
              1 S L84 AND STEINMAN L?/AU
             .5 S L85-L89
L90
              3 S L90 NOT (ZEBRAFISH OR NEMATODE)
L91
              6 S L84 NOT L90
L92
```

FILE 'WPIX' ENTERED AT 16:21:20 ON 01 JUL 2003